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(54) Title: SUBSTRATE TRAPPING PROTEIN TYROSINE PHOSPHATASES

(57) Abstract

Novel protein tyrosine phosphatases in which the invariant aspartate residue is replaced with an alanine residue and which bind to a tyrosine phosphorylated substrate and are catalytically attenuated are described. Also described are methods of identifying tyrosine phosphorylated proteins which complex with the described protein tyrosine phosphatases.

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SUBSTRATE TRAPPING PROTEIN TYROSINE PHOSPHATASES

Background of the Invention

The protein tyrosine phosphatase (PTP) family of enzymes consists of more than 500 structurally diverse proteins which have in common the highly conserved 250 amino acid PTP catalytic domain, but which display considerable variation in their non-catalytic segments (Charbonneau and Tonks, Annu. Rev. Cell Biol. 8:463-493 (1992); Tonks, Semin. Cell Biol. 4:373-453 (1993)). 10 structural diversity presumably reflects the diversity of physiological roles of individual PTP family members, which in certain cases have been demonstrated to have specific functions in growth, development and differentiation (Desai et al., Cell 84:599-609 (1996); Kishihara et al., Cell 74:143-156 (1993); Perkins et al., Cell 70:225-236 (1992); 15 Pingel and Thomas, Cell 58:1055-1065 (1989); Schultz et al., Cell 73:1445-1454 (1993)). Although recent studies have also generated considerable information regarding the structure, expression and regulation of PTPs, the nature of the tyrosine phosphorylated substrates through which the 20 PTPs exert their effects remains to be determined. Studies with a limited number of synthetic phosphopeptide substrates have demonstrated some differences in substrate selectivity of different PTPs (Cho et al., Protein Sci. 2: 977-984 (1993); Dechert et al., Eur. J. Biochem. 231:673-25 681 (1995)), and have indicated preferences for certain amino acid residues at particular positions around the phosphorylated tyrosine residue (Ruzzene et al., Eur. J. Biochem. 211:289-295 (1993); Zhang et al., Biochemistry 30 33:2285-2290 (1994)). This indicates that PTPs display a certain level of substrate selectivity in vitro, although

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the physiological relevance of the substrates used in these studies is unclear.

Summary of the Invention

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As described herein, the substrate specificity of mammalian protein tyrosine phosphatases (PTPs) has been investigated using a novel substrate trapping approach in which mutant or altered forms of the mammalian PTP, also referred to as substrate trapping PTPs, are used to bind (trap) one or more substrates of the PTP. Binding of the substrate trapping PTP with a substrate of the PTP results in the formation of a complex which can be readily observed, and, if desired, isolated, and characterized. The mutant forms of the PTPs have attenuated catalytic activity (lack catalytic activity or have reduced catalytic activity) relative to the wild type PTP but retain the ability to bind tyrosine phosphorylated substrate(s) of the wild type PTP.

The methods of the present invention are specifically exempliefied herein with respect to the phosphatases PTP1B and PTP-PEST; however, it is understood that the invention is not limited to these specific PTPs but is applicable to all members of the PTP family. In order to identify potential substrates of PTP1B and PTP-PEST, mutant (i.e., altered or substrate trapping) forms of PTP1B and PTP-PEST were generated which were catalytically attenuated but retained the ability to bind substrates. These mutant PTPs associated in stable complexes with proteins which were identified by immunoblotting as p210 bcr:abl and p130 cas, respectively. These associations were observed in lysates from several cell lines and in transfected COS cells, indicating that p210 bcr:abl and p130 represent major physiologically relevant substrates for PTP1B and PTP-PEST.

These results provide the first demonstration of PTPs having inherently restricted substrate specificity <u>in vivo</u>.

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The methods used to identify p210 bcr:abl and p130 as specific substrates for PTP1B and PTP-PEST, respectively, are generally applicable to any member of the PTP family, of which approximately 500 members have currently been reported, and can be used to determine the physiological substrates of other members of the PTP family.

One embodiment of the invention relates to novel mutant PTPs in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than These PTPs retain the ability to form a complex with, or bind, their tyrosine phosphorylated substrates, but In one embodiment, the are catalytically attenuated. invention relates to the phosphatase PTP1B in which the 15 invariant aspartate residue at position 181 is replaced with In another embodiment the invention alanine (D181A). relates to the phosphatase PTP-PEST in which the invariant aspartate residue at position 199 is replaced with an alanine (D199A). Another embodiment of the invention 20 relates to a PTP-PEST phosphatase in which the cysteine residue at position 231 is replaced with a serine (C231S). The invention also relates to other mutant or substrate trapping PTPs in which the invariant aspartate residue is replaced with or changed to another amino acid residue, such 25 The invariant aspartate residue can be as alanine. identified in other PTPs by aligning the PTP nucleotide sequence with the nucleotide sequence of a PTP for which the location of the invariant aspartate residue is known.

The invention also relates to a method of identifying a tyrosine phosphorylated substrate of a protein tyrosine phosphatase. According to one embodiment of the present invention, a tyrosine phosphorylated protein of interest is combined with one or more PTP(s) in which the invariant aspartate residue is replaced with an amino acid which does

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not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹), and the presence or absence of a complex between the protein and the PTP(s) is determined. 5 Presence of a complex in the combination indicates that the tyrosine phosphorylated protein is a substrate of the PTP. The PTP DA mutant binds to or complexes with its substrate but does not dephosphorylate it (or does so very slowly), thereby allowing the complex to be observed and, optionally, isolated and identified. In a particular embodiment of the invention, the invariant aspartate is replaced with an alanine residue (a PTP DA mutation or alteration)

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In an alternative embodiment of the present invention, a PTP of interest in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹), is combined with one or more tyrosine phosphorylated proteins, and the presence or absence of a complex between the protein(s) and the PTP is determined. Presence of a complex in the combination indicates that the tyrosine phosphorylated protein is a substrate of the PTP. The PTP DA mutant binds to or complexes with its substrate but does not dephosphorylate it (or does so very slowly), thereby allowing the complex to be observed, and, optionally, isolated and identified. In one embodiment of the invention, the invariant aspartate residue is replaced with an alanine residue (a PTP DA mutation or alteration)

The present invention also relates to a method of identifying a tyrosine phosphorylated substrate of a protein tyrosine phosphatase wherein more than one tyrosine phosphorylated protein of interest is combined with more than one PTP of interest in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which

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results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹) (e.g., the invariant aspartate is replaced with an alanine residue). Complexes formed in the combination can be isolated and the component PTP and substrate can be identified.

The invention also pertains to a method of reducing the activity of a tyrosine phosphorylated protein, comprising administering to a mammal a PTP in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹) (e.g., the invariant aspartate is replaced with an alanine residue) and which forms a complex with the tyrosine phosphorylated protein. The PTP mutant binds to the phosphorylated protein without dephosphorylating it, thereby inhibiting the activity of the protein and reducing its downstream effects.

For example, the invention relates to a method of reducing the transforming effects of oncogenes associated with pl30 cas, a substrate of PTP-PEST, comprising 20 administering to a mammal wild type PTP-PEST or PTP-PEST in which the invariant aspartate residue is replaced with an alanine residue. Wild type PTP-PEST binds and dephosphorylates p130 cas, thereby negatively regulating its downstream effects. DA mutants of PTP-PEST bind but do not 25 dephosphorylate p130 cas (or dephosphorylate it at a reduced rate); the substrate is thus tied up in the complex with the substrate trapping form of PTP-PEST and cannot exert its downstream effects. Similarly, the invention relates to a method of reducing the formation of signalling complexes 30 associated with p130 cas, particularly those signalling complexes which induce mitogenic pathways, comprising administering to a mammal wild type PTP-PEST or PTP-PEST in which the invariant aspartate residue is replaced with an alanine residue.

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The present invention also relates to assays for identifying agents which alter, e.g., enhance or inhibit, the interaction between a PTP and its phosphorylated substrate. Agents identified by these assays can be agonists (e.g., agents which enhance or increase the activity of the PTP) or antagonists (e.g., agents which inhibit or decrease the activity of the PTP) of PTP activity. The agent may be an endogenous physiological substance or may be a natural or synthetic drug, including small organic molecules.

For example, the tyrosine phosphorylated substrate of a PTP can be identified by the methods described herein. enzymatic activity assay utilizing the wild type PTP can be carried out in the presence of an agent to be tested, and 15 the resulting amount of enzyme activity can be compared with the amount of enzyme activity in the absence of the agent to be tested. A decrease in the enzymatic activity in the presence of the agent to be tested indicates that the agent inhibits the interaction between the PTP and its substrate. 20 Conversely, an increase in the enzymatic activity in the presence of the agent to be tested indicates that the agent enhances the interaction between the PTP and its substrate.

Alternatively, a competitive binding assay can be carried out utilizing the mutant PTP in the presence of an agent to be tested, and the resulting extent of binding of the mutant PTP to its substrate can be compared with the extent of binding in the absence of the agent to be tested. A decrease in the extent of binding in the presence of the agent to be tested indicates that the agent inhibits the 30 interaction between the PTP and its substrate. Conversely, an increase in the extent of binding in the presence of the agent to be tested indicates that the agent enhances the interaction between the PTP and its substrate.

Thus, the compositions and methods described herein are useful in identifying the tyrosine phosphorylated 35

substrates of members of the PTP family of phosphatases, as well as in regulating the activity of identified substrates. The compositions and methods described herein are also useful for identifying tyrosine phosphorylated proteins which are related to a particular disease or disorder, and to methods of screening for modulators which enhance or inhibit the PTP/substrate interaction for use in therapeutic applications.

Brief Description of the Drawings

Figures 1A and 1B show a multiple sequence alignment 10 of the catalytic domains of PTPs. In Figure 1A, cytosolic eukaryotic PTPs and domain 1 of RPTPs are combined into one group, domains 2 of RPTPs are in a second group, and the Yersinia PTP is in a third. Invariant residues shared among all three groups are shown in red. Invariant and highly conserved residues within a group are shown in blue and green, respectively. Within the Yersinia PTP sequence, residue that are either invariant or highly conserved between the cytosolic and RPTP domain sequences are colored blue and green, respectively. The position of residues of PTP1B that interact with the peptide are indicated with a red arrow, and the residue numbering at the bottom of the alignment corresponds to that for PTP1B. Figure 1B is a black and white photocopy of Figure 1A in which the colored areas are indicated with labeled arrows.

Figure 2 shows the Vmax and Km of various PTP1B mutants toward RCML.

Detailed Description of the Invention

The PTP family of enzymes contains a common evolutionarily conserved segment of approximately 250 amino acids known as the PTP catalytic domain. Within this

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conserved domain is a unique signature sequence motif, [I/V] HCXAGXXR[S/T]G, that is invariant among all PTPs. cysteine residue in this motif is invariant in members of the family and is known to be essential for catalysis. 5 functions as a nucleophile to attack the phosphate moiety of the incoming substrate. If the cysteine residue is altered by site-directed mutagenesis to serine (CS mutants) or alanine (CA mutants), the resulting PTP is catalytically attenuated but retains the ability to complex with, or bind, its substrate, at least in vitro. These results have been 10 confirmed relative to MKP-1, a member of the PTP family (Sun et al., Cell 75:487-493 (1993)), as well as other PTPs. However, although these CS mutants can in general bind effectively to phosphotyrosyl substrates in vitro, in many cases such complexes cannot be isolated in vivo. Thus, the 15 CS mutants are limited in their applicability and cannot be used to isolate all combinations of PTPs and substrates. The crystal structures of PTP1B alone (Barford, et al., Science 263:1397-1404 (1994)) and in a complex with a phosphotyrosine-containing peptide (Jia et al., Science 20 268:1754-1758 (1995)) were recently determined. structures indicated twenty seven invariant residues (Barford et al., 1994), one of which is an aspartate This aspartate residue is invariant across the 25 catalytic domains of PTP family members. That is, if the amino acid sequences of the PTP family members are aligned, the aspartate residue is present in each PTP at a corresponding location, although the position numbers may be different due to the shifts required to maximize alignment (see the Figure (from Barford et al., Nature Struc. Biol. 30 2:1043-1053 (1995)) for an alignment of various PTP sequences). Sequences for which the alignment has not yet been published can readily be aligned with other known PTP sequences, e.g., utilizing available computer software such

Thus, mutant PTPs other than those specifically described herein can readily be made by aligning the amino acid sequence of the PTP catalytic domain with those described herein, identifying the invariant aspartate residue, and changing the residue by site-directed mutagenesis. Although the specific examples of PTP mutants described herein are aspartate to alanine mutants (DA mutants), it is understood that the invention is not limited to changes of aspartate to alanine. The invariant aspartate residue can be changed, e.g., by site-directed mutagenesis, 10 to any amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min 1). For example, the invariant aspartate residue can be changed or mutated to an alanine, valine, leucine, isoleucine, proline, phenylalanine, tryptophan, methionine, glycine, serine, threonine, cysteine, tyrosine, asparagine, glutamine, lysine, arginine or histidine.

As described herein, pervanadate-treated cells were used as an abundant source of tyrosine phosphorylated 20 proteins to investigate the substrate specificity of PTP-PEST. PTP-PEST is an 88 kDa cytosolic PTP (Charest et al., Biochem. J. 308:425-432 (1995); den Hertog et al., Biochem. Biophys. Res. Commun. 184:1241-1249 (1992); Takekawa et al., Biochem. Biophys. Res. Commun. 189:1223-1230 (1992); Yang et al., J. Biol. Chem. 268:6622-6628 (1993); Yang et al., J. Biol. Chem. 268:17650 (1993)) which is expressed ubiquitously in mammalian tissues (Yi et al., Blood 78: 2222-2228 (1991)), and which exhibits high specific activity when assayed in vitro using artificial tyrosine 30 phosphorylated substrates (Garton and Tonks, EMBO J. 13:3763-3771 (1994)). It has previously been demonstrated that PTP-PEST is subject to regulation via phosphorylation of Ser39 in vitro and in vivo. This modification is catalyzed by both protein kinase C (PKC) and protein kinase 35

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A (PKA), and results in reduced enzyme activity as a consequence of an increase in the Km of the dephosphorylation reaction (Garton and Tonks, EMBO J. 13:3763-3771 (1994)). It appears likely that further 5 regulatory mechanisms exist for PTP-PEST, since this enzyme would be expected to exert a considerable negative influence on the tyrosine phosphorylation state of cytosolic substrates of tyrosine kinases. One possibility is that this influence could be limited by the substrate specificity of PTP-PEST.

The substrate specificity of PTP1B was investigated utilizing the same methods outlined for PTP-PEST, with the exception that the cells were not treated with pervanadate. A combination of in vitro dephosphorylation and substrate trapping experiments were used to study the substrate interactions of PTP1B and PTP-PEST. The substrate trapping methods outlined herein are generally applicable to any PTP by virtue of the shared invariant aspartate residue, and should therefore prove useful in delineating the substrate preference of other PTP family members. In particular, the use of mutant, catalytically impaired PTPs to trap, and thereby isolate, potential substrates will greatly facilitate the identification of physiologically important substrates for individual PTPs, leading to improved understanding of the roles of these enzymes in regulation of cellular processes.

One embodiment of the invention relates to novel PTPs in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min-1). PTPs retain the ability to form a complex with, or bind, their tyrosine phosphorylated substrates but are catalytically attenuated. As defined herein, "attenuated" activity is intended to mean that the phosphatase retains a

similar Km to that of the wild type phosphatase but has a Vmax which is reduced by a factor of at least 10⁴ relative to the wild type enzyme. This includes catalytic activity which is either reduced or abolished relative to the wild type PTP. For example, the invariant aspartate residue can be changed or mutated to an alanine, valine, leucine, isoleucine, proline, phenylalanine, tryptophan, methionine, glycine, serine, threonine, cysteine, tyrosine, asparagine, glutamine, lysine, arginine or histidine.

The novel PTPs described herein, in which the 10 invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹), can also comprise other mutations, particularly those which assist in stabilizing 15 the PTP/substrate complex. For example, a mutation of the [serine/threonine] residue in the signature motif to an alanine residue changes the rate-determining step of the dephosphorylation reaction from the formation of the 20 transition state to the break down of the transition state, thereby stabilizing the PTP/substrate complex. Such mutations may be valuably combined with the replacement of the invariant aspartate residue, particularly assisting in stabilizing the complex and facilitating the observation and isolation of the complex. 25

ptps suitable for use in the invention include any PTP
which has an invariant aspartate residue in a corresponding
position. As defined herein, a phosphatase is a member of
the PTP family if it contains the signature motif

[I/V]HCXAGXXR[S/T]G. Dual specificity PTPs, i.e., PTPs
which dephosphorylate both phosphorylated tyrosine and
phosphorylated serine or threonine, are also suitable for
use in the invention. Appropriate PTPs include, but are not
limited to, PTP1B, PTP-PEST, PTP, MKP-1, DEP-1, PTPμ,

PTPX1, PTPX10 and PTPH1.

In one embodiment, the invention relates to the phosphatase PTP1B in which the aspartate residue at position 181 is replaced with alanine (D181A). In another embodiment the invention relates to the phosphatase PTP-PEST in which the invariant aspartate residue at position 199 is replaced with an alanine (D199A). Another embodiment of the invention relates to a PTP-PEST phosphatase in which the cysteine residue at position 231 is replaced with a serine (C231S).

10 The invention also relates to a method of identifying a tyrosine phosphorylated protein which is a substrate of a particular protein tyrosine phosphatase. According to one embodiment of the present invention, a tyrosine phosphorylated protein of interest is combined with at least 15 one PTP in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹) (e.g., an alanine residue), and the presence or absence of a complex between the protein and the PTP is 20 determined. Presence of a complex in the combination indicates that the tyrosine phosphorylated protein is a substrate of the PTP. The PTP DA mutant (substrate trapping mutant) binds to or complexes with its substrate but does 25 not dephosphorylate it (or does so very slowly), thereby allowing the complex to be isolated and identified.

The phosphorylated protein/PTP complex may be isolated by conventional isolation techniques as described in U.S. Patent No. 5,352,660 to Pawson, including salting out, chromatography, electrophoresis, gel filtration, fractionation, absorption, polyacrylamide gel electrophoresis, agglutination, or combinations thereof. Furthermore, to facilitate the determination of the presence of the protein/PTP complex, antibodies against the PTP or the phosphorylated protein can be used, as well as labelled

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PTPs and/or labelled phosphorylated substrates. The PTP or phosphorylated protein can be labelled with various enzymes, fluorescent materials, luminescent materials and radioactive materials. Examples of suitable enzymes include, but are not limited to, horseradish peroxidase, biotin, alkaline phosphatase, β-galactosidase and acetylcholinesterase. Examples of suitable fluorescent materials include, but are not limited to, umbelliferone, fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine

10 fluorescein, dansyl chloride and phycoerythrin. Appropriate luminescent materials include luminol, and suitable radioactive material include radioactive phosphorous ³²P, iodine I¹²⁵, I¹³¹ or tritium.

Alternatively, the invention pertains to a method of identifying a tyrosine phosphorylated protein which is a 15 substrate of a PTP, comprising combining a PTP of interest in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹) (e.g., an 20 alanine residue), with at least one tyrosine phosphorylated protein, thereby producing a combination; and determining the presence or absence of a complex in the combination, wherein presence of a complex in the combination between a tyrosine phosphorylated protein and the PTP indicates that 25 the tyrosine phosphorylated protein is a substrate of the

The substrate trapping PTPs of the present invention can also be used in place of wild type PTPs to screen phosphotyrosyl peptide libraries for peptides which bind to the PTP as described in Songyang et al. (Nature 373:536-539 (1995); Cell 72:767-778 (1993)). Peptides identified from such peptide libraries can then be assessed to determine whether tyrosine phosphorylated proteins containing these peptides exist in nature.

Any tyrosine phosphorylated protein is suitable as a potential substrate in the present invention. Tyrosine phosphorylated proteins are well known in the art. Specific examples of appropriate substrates include, without limitation, pl30 cas, the EGF receptor, p210 bcr:abl, MAP kinase and the insulin receptor. Of particular interest are tyrosine phosphorylated proteins which have been implicated in a mammalian disease or disorder.

The invention also pertains to a method of reducing the activity of a tyrosine phosphorylated protein, 10 comprising administering to a mammal a PTP in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹) (e.g., an alanine residue) 15 and which forms a complex with the tyrosine phosphorylated protein. The PTP DA mutant binds to the phosphorylated protein without dephosphorylating it (or causing dephsophorylation at a greatly reduced rate), thereby inhibiting the activity of the protein and reducing its 20 downstream effects. As used herein, "reducing" includes both reduction and complete abolishment, e.q., of one or more activities or functions of the phosphorylated protein.

For example, the invention relates to a method of

25 reducing the transforming effects of oncogenes associated with p130 cas, a substrate of PTP-PEST, comprising administering to a mammal wild type PTP-PEST or PTP-PEST in which the invariant aspartate residue is replaced with an alanine residue. Wild type PTP-PEST binds and

30 dephosphorylates p130 cas, thereby negatively regulating its downstream effects. DA mutants of PTP-PEST bind but do not dephosphorylate p130 (or do so at a greatly reduced rate); the substrate is thus tied up in the complex with the substrate trapping form of PTP-PEST and cannot exert its

35 downstream effects. Similarly, the invention relates to a

method of reducing the formation of signalling complexes associated with $p130^{\text{Cas}}$, particularly those signalling complexes which induce mitogenic pathways, comprising administering to a mammal wild type PTP-PEST or PTP-PEST in which the invariant aspartate residue is replaced with an alanine residue. The PTP binds to and/or dephosphorylates $p130^{\text{Cas}}$, thereby negatively regulating the downstream effects of $p130^{\text{Cas}}$ and reducing the formation of signalling complexes associated with $p130^{\text{Cas}}$.

The substrate trapping mutant PTPs of the present 10 invention can be used in virtually any application in place of, or in addition to, a corresponding wild type PTP. advantages of such a utility lie in the ability of the mutant PTP to mimic the function of the wild type enzyme, e.g., to decrease the activity of its tyrosine 15 phosphorylated substrate, without inducing the harmful cytotoxic effects commonly observed with administration or overexpression of the wild type PTP. Thus, the invention also pertains to a method of reducing the cytotoxic effects associated with administration or overexpression of wild 20 type PTPs. For example, CS mutants of MKP-1 have been shown to have the same functional effect as wild type MKP-1 without induction of potentially harmful side effects. Thus, PTPs described herein, in which the invariant aspartate residue is replaced with an amino acid which does 25 not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min⁻¹) (e.g., an alanine residue), can be used in many applications in place of the corresponding wild type enzyme. As used herein, a "corresponding" enzyme is one which is the same as the mutant PTP (e.g., PTP-PEST and PTP-PEST D199A) or one which is different from the mutant PTP but recognizes the same substrate as the mutant PTP.

The mutant PTPs described herein can also be used therapeutically to reduce the activity of a tyrosine phosphorylated protein, such as by a gene therapy method in which the mutant PTP described herein, or a functional 5 portion thereof which retains the ability to bind to its tyrosine phosphorylated substrate, is introduced into a subject and in whom the mutant PTP is expressed. PTP replaces, either partially or totally, the wild type enzyme which is normally produced or competes with the wild 10 type PTP for binding to the substrate. For example, a specific tyrosine phosphorylated protein can be identified which is implicated in a particular disease or disorder (such as a protein tyrosine kinase). At least one PTP which acts to dephosphorylate the selected tyrosine phosphorylated protein of the present invention can be identified by the 15 methods described herein. The wild type or mutant form of the PTP can be administered to a subject in need of treatment in order to tie up or bind the tyrosine phosphorylated substrate, thereby inhibiting or reducing the function of the phosphorylated protein. In a preferred 20 embodiment, the mutant PTP is administered in place of the wild type enzyme in order to reduce the cytotoxic effects associated with overexpression of the wild type enzyme. Procedures for gene therapy are known in the art (see U.S. Patent No. 5,399,346 to Anderson et al.) and can be modified by methods known in the art to appropriately express the specific mutant and wild type PTPs of the present invention.

The present invention also relates to assays for identifying agents which alter, e.g., enhance or inhibit, 30 the interaction between a PTP and its phosphorylated substrate. Agents identified by these assays can be agonists (e.g., agents which enhance or increase the activity of the PTP) or antagonists (e.g., agents which inhibit or decrease the activity of the PTP) of PTP activity. The agent may be an endogenous physiological

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substance or may be a natural or synthetic drug, including small organic molecules.

For example, the tyrosine phosphorylated substrate of a PTP can be identified by the methods described herein. 5 enzymatic activity assay utilizing the wild type PTP can be carried out in the presence of an agent to be tested, and the resulting amount of enzyme activity can be compared with the amount of enzyme activity in the absence of the agent to be tested. Enzymatic activity assays are known in the art; 10 for example, assays of PTP activity using a tyrosine phosphorylated ³²P-labelled substrate are described in Flint et al. (EMBO J. 12:1937-1946 (1993)). A decrease in the enzymatic activity in the presence of the agent to be tested indicates that the agent inhibits the interaction between the PTP and its substrate. Conversely, an increase 15 in the enzymatic activity in the presence of the agent to be tested indicates that the agent enhances the interaction between the PTP and its substrate.

Alternatively, a competitive binding assay can be carried out utilizing the mutant PTP in the presence of an agent to be tested, and the resulting extent of binding of the mutant PTP to its substrate can be compared with the extent of binding in the absence of the agent to be tested. Competitive binding assays are known in the art; for example, U.S. Patent No. 5,352,660 to Pawson describes methods suitable for use in this invention. A decrease in the extent of binding in the presence of the agent to be tested indicates that the agent inhibits the interaction between the PTP and its substrate. Conversely, an increase in the extent of binding in the presence of the agent to be tested indicates that the agent enhances the interaction between the PTP and its substrate.

According to the present invention, tyrosine phosphorylated peptides identified with mutant PTPs from peptide libraries by the methods of Songyang et al. (Nature

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373:536-539 (1995); Cell 72:767-778 (1993)) can be used herein in place of the complete tyrosine phosphorylated protein in competitive binding assays.

The present invention also pertains to pharmaceutical 5 compositions comprising a PTP in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute (less than 1 min.1) (e.g., an alanine residue). For instance, the PTP of the present invention can be formulated with a physiologically acceptable medium to prepare a pharmaceutical composition. The particular physiological medium may include, but is not limited to, water, buffered saline, polyols (e.g., glycerol, propylene glycol, liquid 15 polyethylene glycol) and dextrose solutions. The optimum concentration of the active ingredient(s) in the chosen medium can be determined empirically, according to procedures well known to medicinal chemists, and will depend on the ultimate pharmaceutical formulation desired. 20 of introduction of exogenous PTPs at the site of treatment include, but are not limited to, intradermal, intramuscular, intraperitoneal, intravenous, subcutaneous, oral and intranasal. Other suitable methods of introduction can also include rechargeable or biodegradable devices and slow release polymeric devices. The pharmaceutical compositions 25 of this invention can also be administered as part of a combinatorial therapy with other agents.

The following Examples are offered for the purpose of illustrating the present invention and are not to be construed to limit the scope of this invention. The teachings of all references cited herein are hereby incorporated by reference in their entirety.

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EXAMPLES

Materials and Methods

The following is a description of the materials and methods used in the work described herein.

5 <u>Generation, Expression and Purification of Mutant PTP</u> <u>Proteins</u>

Point mutations within the catalytic domains of PTP-PEST (D199A, C231S) and PTP1B (D181A, C215S) were introduced by site-directed mutagenesis using the Muta-Gene in vitro mutagenesis kit (Bio-Rad, Richmond, CA). Regions containing the required point mutation were then exchanged with the wild type sequences within appropriate expression vectors, and the replaced mutant regions were sequenced in their entirety to verify the absence of additional mutations.

Full length PTP-PEST proteins (wild type and mutant 15 proteins, containing either Asp199 to Ala or Cys231 to Ser mutations) and the wild type PTP-PEST catalytic domain (amino acids 1-305) were expressed in Sf9 cells using recombinant baculovirus (BaculoGold , Pharmingen, San 20 Diego, CA), and purified as described in Garton and Tonks (<u>EMBO J. 13</u>:3763-3771 (1994)). Truncated forms of wild type and mutant PTP-PEST proteins, comprising amino acid residues 1-305 of PTP-PEST were also expressed in E. coli as GST fusion proteins following subcloning of PTP-PEST DNA in-25 frame downstream of GST in pGEX vectors (Pharmacia Biotech Inc., Uppsala, Sweden). Twenty-five ml of E. coli transformed with the appropriate vector were grown to log phase $(OD_{600}$ approximately 0.5). Fusion protein expression was then induced by addition of 0.2 mM isopropyl-1-thio-b-Dgalactopyranoside, and the cells were grown for 2-4 hours at 30 Cells were harvested by centrifugation, incubated with 50 mg/ml lysozyme in 3 ml buffer containing 50 mM Tris-HCl, pH 7.4, 5mM EDTA, 1 mM PMSF, 1 mM benzamidine, 5 mg/ml leupeptin, 5 mg/ml aprotinin, 0.1 % Triton X-100 and 150 mM

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NaCl, then lysed by sonication (3 \times 10s). Following removal of insoluble material by centrifugation (20 minutes at 300,000 \times g), fusion proteins were isolated by incubation for 30 min at 4°C with 100 ml glutathione-Sepharose beads 5 (Pharmacia Biotech Inc., Uppsala, Sweden), and the beads were then collected by centrifugation and washed three times with buffer A (20 mM Tris-HCl, pH 7.4, 1 mM EDTA, 1 mM benzamidine, 1 mg/ml leupeptin, 1 mg/ml aprotinin, 10 % glycerol, 1 % Triton X-100 and 100 mM NaCl). This procedure yielded essentially homogeneous fusion protein at a 10 concentration of 1 mg protein/ml glutathione-Sepharose beads. PTP1B proteins (wild type and mutant forms) comprising amino acids 1-321 were expressed in E.coli and purified to homogeneity as described in Barford et al. (J. 15 Mol. Biol. 239:726-730 (1994)).

<u>Cell Culture, Transfection, Preparation of Lysates and Fractionation</u>

HeLa and COS cells were grown in Dulbecco's modified Eagle's medium (DMEM), containing 5% fetal bovine serum (FBS); Wi38, C2C12 and MvLu cells were grown in DMEM 20 containing 10% FBS; 293 cells were grown in DMEM containing 10% calf serum; MCF10A cells were grown in 50% DMEM, 50% Ham's F-12 containing 5% horse serum, 20 ng/ml epidermal growth factor, 10 mg/ml insulin, 0.5 mg/ml hydrocortisone and 0.25 mg/ml fungizone. All media also contained 25 penicillin and streptomycin at 100 U/ml and 100 mg/ml, respectively, and all cells were grown at 37°C. Calcium phosphate-mediated transfection was used to introduce cDNA encoding wild type and mutant PTP-PEST proteins into COS cells. These were encoded by PTP-PEST cDNA subcloned into 30 the plasmid PMT2, from which expression was driven by an adenovirus major late promoter; 20 mg DNA was used for transfection of each 10 cm plate of cells. The level of expression of PTP-PEST constructs was similar in all cases.

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Prior to cell lysis, 70-90% confluent cultures of cells were treated for 30 minutes with 0.1 mM pervanadate (20 ml of a fresh solution containing 50 mM sodium metavanadate (NaVO₃) and 50 mM H₂O₂ were added to 10 ml medium). Treatment of cells with ${\rm H}_2{\rm O}_2$ and vanadate leads to a synergistic increase in phosphotyrosine levels, presumably due to inhibition of intracellular PTPs by vanadate. The synergism between H2O2 and vanadate has previously been suggested to result from improved accumulation of the resultant oxidized vanadate 10 (pervanadate) within the cells when compared to vanadate itself (Heffetz et al., J. Biol. Chem. 265:2896-2902 (1990)). It is important to note that during the preparation of cell lysates, dilution occurs such that the 15 inhibitory effect of vanadate on PTP action is lost. Pervanadate treatment resulted in the appearance of at least 50 prominent phosphotyrosine protein bands in all cell types, whereas untreated cells contained virtually undetectable levels of phosphotyrosine (data not shown).

Cells were lysed in Buffer A containing 5 mM iodoacetic acid, which was included in order to inhibit irreversibly cellular PTPs. Following incubation at 4°C for 30 minutes, 10 mM DTT was added to inactivate any unreacted iodoacetic acid. Insoluble material was then removed by centrifugation for 20 minutes at 300,000 x g. The resultant lysates were stable with regard to their phosphotyrosine content during long term (several months) storage at -70°C and during prolonged (at least 20 hours) incubation at 4°C, in the absence of exogenous added PTPs.

Pervanadate-treated HeLa cell lysate was fractionated by anion exchange chromatography using a Mono Q FPLC column (Pharmacia). The sample (50 mg total protein at 3 mg/ml in buffer A) was diluted in three volumes of buffer B (20 mM tris-HCl, pH 7.4, 1 mM EDTA, 1 mM benzamidine, 1 mg/ml leupeptin, 1 mg/ml aprotinin and 0.1% Triton X-100) prior to

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loading. Proteins were eluted at a flow rate of 1 ml/min with a linear gradient of 0-0.5 M NaCl in buffer B over 20 fractions (1 ml fraction volume), followed by a second gradient of 0.5-1.0 M NaCl in buffer B over 5 fractions. Phosphotyrosine-containing proteins were detected within fractions 7-21 according to anti-phosphotyrosine immunoblotting. The same procedures were followed for PTP1B, with the exception that the cells were not treated with pervanadate.

10 <u>Dephosphorylation Reactions</u>

Lysates of pervanadate-treated HeLa cells (1-2 mg protein/ml) containing tyrosine phosphorylated proteins were incubated on ice in the absence or presence of purified active PTPs at a concentration of 2 nM. Dephosphorylation

15 was terminated by the removal of aliquots (30 mg protein) into SDS-PAGE sample buffer, and the extent of dephosphorylation was determined by immunoblotting using the monoclonal antibody G104. Assays of PTP activity using tyrosine phosphorylated TP-labelled RCM-lysozyme as substrate were performed as described in Flint et al. (EMBO J. 12:1937-1946 (1993)).

Antibodies and Immunoblotting

The PTP-PEST monoclonal antibody AG25 was raised against baculovirus-expressed purified full-length PTP-PEST.

The anti-phosphotyrosine monoclonal antibody G104 was generated using as antigen phosphotyrosine, alanine and glycine, in a 1:1:1 ratio, polymerized in the presence of keyhole limpet hemocyanin with 1-ethyl-3-(3'-dimethylaminopropyl)carbodiimide, a method originally described in Kamps and Sefton (Oncogene 2:305-315 (1988)). p130 monoclonal antibody was from Transduction Laboratories (Lexington, Ky). Monoclonal antibody FG6 against PTP1B was provided by Dr David Hill (Calbiochem

Oncogene Research Products, Cambridge, MA). Visualization of proteins by immunoblotting was achieved by enhanced chemiluminescence (ECL) using HRP-conjugated secondary antibodies (Amersham Life Science Inc., Arlington Heights, Il) and the SuperSignal CL-HRP substrate system (Pierce, Rockford, Il).

Immunoprecipitation and Substrate Trapping

Immunoprecipitation of PTP-PEST from transfected COS cells was performed following covalent coupling of monoclonal antibody AG25 to protein A-Sepharose beads 10 (Pharmacia Biotech Inc., Uppsala, Sweden) using the chemical cross-linking agent dimethyl pimelimidate (Schneider et al., <u>J. Biol. Chem.</u> <u>257</u>:10766-10769 (1982)). Antibody was first bound to protein A-Sepharose at a concentration of 1 mg/ml bead volume, and unbound material was then removed by three 15 washes with 0.2 M sodium borate, pH 9. Covalent coupling was achieved by incubation at room temperature for 30 minutes in the presence of 20 mM dimethyl pimelimidate in 0.2 M sodium borate, pH 9. The beads were then incubated for 1 hour with an excess of 0.2 M ethanolamine, pH 8, to 20 block any unreacted cross-linker, and washed three times with PBS prior to storage at 4°C. Ten ml of AG25 beads were used to precipitate transfected PTP-PEST from lysates containing approximately 0.375 mg protein.

Substrate trapping was performed using various PTP affinity matrices. The full-length PTP-PEST matrix utilized covalent coupled AG25-protein A-Sepharose beads to which purified baculovirus-expressed PTP-PEST protein was bound. Aliquots (10ml) of AG25 beads were incubated for 2 hours at 4°C in 100 ml buffer A in the presence of 5 mg of purified PTP-PEST (wild type or mutant forms); unbound PTP-PEST was then removed by washing three times with 1 ml buffer A. The resultant PTP-PEST-AG25-protein A-Sepharose beads contained approximately 2 mg of PTP-PEST per 10 ml aliquot. Substrate

trapping was also carried out with glutathione-Sepharose beads bound to bacterially-expressed GST fusion proteins containing the catalytic domain of PTP-PEST.

PTP1B was also used in substrate trapping experiments. In this case, the monoclonal antibody FG6 was precoupled to protein A-Sepharose in the absence of cross-linker (2 mg antibody/10 ml beads), then purified PTP1B proteins were added in excess and incubated at 4°C for 2 hours. Following removal of unbound PTP1B, 10 ml beads contained approximately 2 mg PTP1B.

Pervanadate-treated cell lysates, or column fractions, were used as a source of phosphotyrosine-containing proteins for substrate trapping experiments. In general, lysates containing 0.25-0.5 mg protein in 0.5 ml buffer A (including 5 mM iodoacetic acid, 10 mM DTT) were incubated at 4°C for 2 hours in the presence of 10 ml of affinity matrix containing approximately 2 mg of the appropriate PTP protein. Unbound proteins were then removed from the samples by washing three times with 1 ml buffer A, and bound material was collected by addition of 50 ml SDS-PAGE sample buffer followed by heating at 95°C for 5 minutes; proteins bound to the beads were then analyzed by SDS-PAGE followed by immunoblotting.

Results

The following details the results of the work described herein carried out as described above.

PTP1B and p210 bcr:abl

Chronic myelogenous leukemia (CML) is a clonal disorder of the haematopoietic stem cell that is characterized by the Philadelphia chromosome, in which the c-Abl proto-oncogene on chromosome 9, encoding a PTK, becomes linked to the bcr gene on chromosome 22. This results in the generation of a bcr:able fusion protein, p210 bcr:abl, in which the PTK activity is enhanced relative to

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that of c-Abl. Current data indicates that this cytogenetic abnormality is the primary and sole causative event in CML. Expression of p210 bcr:abl produces abnormal patterns of tyrosine phosphorylation that result in the aberrant maturation of the haematopoietic stem cell that is characteristic of CML.

Expression of PTP1B mRNA and protein is enhanced as a consequence of p210 bcr:abl expression in Rat1, Mo7 and BaF3 cells. Changes in PTP1B activity, which were commensurate with the change in enzyme protein, were also observed. These changes are specific for PTP1B and are not seen in closely related homologue (65% identity) TC-PTP or in other tested PTPs, including SHP-1, SHP-2 and PTP-PEST. The increase in expression of PTP1B was also observed in Ph+ B-lymphoid cells derived from a CML patient relative to Ph-cells from the same patient.

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The changes in PTP1B levels were induced specifically by p210 bcr:abl and were not seen in cells expressing other PTKs including v-abl, v-src or other oncoproteins such as 20 The PTK activity of p210 bcr:abl was essential for the increase in expression of PTP1B, since expression of an inactive lysine to arginine mutant form of p210 bcr:abl in Rat1 cells did not alter PTP1B levels. The increase in PTP1B levels is a rapid response to induction of p210 25 bcr:abl. When BaF3 cells expressing a temperature-sensitive mutant form of p210 bcr:abl were shifted to the permissive temperature for the PTK, PTP1B levels were observed to increase within 12-24 hours coincident with the appearance of the active form of the PTK. These data indicate that the alteration PTP1B levels is a relatively rapid response to 30 the appearance of p210 bcr:abl, rather than a long-term adaptive response of the cells.

In transient cotransfection experiments in COS cells,
PTP1B dephosphorylates p210 bcr:abl but not v-abl. When the
PTP1B D181A mutant was expressed as a GST fusion protein,

purified and incubated with lysates of Mo7-p210 cells (which overexpress p210 bcr:abl), a complex of the mutant PTP and p210 bcr:abl was isolated. In contrast, tyrosine phosphorylated c-abl, which was also present in the lysates, did not bind to the mutant PTP. The interaction between PTP1B D181A and p210 bcr:abl was blocked by vanadate, suggesting that the interaction involved the active site of the PTP.

Following transient coexpression in COS cells, PTP1B 10 D181A formed a complex with p210 bcr:abl. Preliminary data indicate that the Y177F mutant form of p210 bcr:abl did not interact with PTP1b D181A, suggesting that this tyrosine residue is a component of the binding site in the PTK. tyrosine residue in p210 bcr:abl is phosphorylated in vivo and has been demonstrated to serve as a docking site for 15 GRB2. Direct interaction of the pTyr in p210 bcr:abl and the SH2 domain of GRB2 is essential for the transforming activity of the PTK. Interaction of PTP1B D181A with p210 bcr:abl interferes with the association of the PTK with GRB2. Taken together, these data suggest that p210 bcr:abl 20 is a physiological substrate of PTP1B and that PTP1B may function as an antagonist of the oncoprotein PTK in vivo. The Vmax, Km and Kcat of 37 kDa PTP1B mutants toward RCML are shown in Figure 2.

25 PTP1B and the EGF Receptor

Expression of PTP1B D181A in COS cells leads to enhanced phosphorylation of tyrosyl residues in a 180 kDa protein and in proteins of 120 and 70 kDa. When a GST-PTP1B D181A fusion protein is expressed in COS cells and precipitated on Glutathione-Sepharose, the 180 kDa, and smaller quantities of p120 and p70, were coprecipitated. The p180 protein was identified as the epidermal growth factor (EGF) receptor by immunoblotting. The identity of

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the pl20 and p70 proteins is unclear; however, the latter is not src, p62 or paxillin.

Expression of PTP1B D181A in COS cells induces tyrosine phosphorylation of the EGF receptor in the absence of its ligand, EGF, indicating that the mutant PTP is exerting its effects in the intact cell and not post-lysis. The equivalent D199A PTP-PEST mutant does not interact with the EGF receptor, indicating the specificity of this substrate interaction.

Autophosphorylation of the EGF receptor is required for the interaction with PTP1B D181A. Mutants of the receptor that are either kinase-dead or in which the autophosphorylation sites have been deleted do not interact with PTP1B D181A. In v-src-expressing cells, a plethora of tyrosine phosphorylated proteins were observed, but phosphorylation of the EGF receptor was not detected. Under these conditions, PTP1B D181A bound predominantly to a 70 kDa tyrosine phosphorylated protein.

As a result of this work, it appears that PTP1B can modulate EGF-induced signalling pathways, perhaps including the pathways of many diseases, including breast cancer.

Preferential Dephosphorylation of a 130 kDa Phosphotyrosine-Containing Protein by PTP-PEST

In order to investigate the substrate specificity of

PTP-PEST in vitro, aliquots of pervanadate-treated HeLa cell
lysates were incubated on ice, yielding 50-100 distinct
phosphotyrosine-containing proteins as judged by
immunoblotting of the cell lysate using the monoclonal antiphosphotyrosine antibody G104. Purified full-length PTP
PEST (expressed in Sf9 cells using recombinant baculovirus),
PTP-PEST catalytic domain, or PTP1B catalytic domain (37 kDa
form) was then added to the lysate, and aliquots were
removed at various time points for analysis by SDS-PAGE
followed by anti-phosphotyrosine immunoblotting.

Surprisingly, a prominent 130 kDa phosphotyrosine band (p130) was selectively dephosphorylated by PTP-PEST within 10 minutes, whereas the intensity of all the other bands was essentially unchanged even after 60 minutes of incubation with PTP-PEST. Long incubations with higher concentrations of PTP-PEST (greater than 100-fold) resulted in the complete removal of all phosphotyrosine bands from the lysate. However, under all conditions tested, p130 was found to be dephosphorylated more rapidly than all other bands present.

The selective dephosphorylation of p130 by PTP-PEST was also observed using a truncated form of the phosphatase (amino acid residues 1-305) which essentially contains only the catalytic domain of the enzyme. This result suggests that the striking substrate preference displayed by PTP-PEST in this analysis is an inherent property of the phosphatase catalytic domain, whereas the C-terminal 500 amino acid residues have little discernible effect on the substrate specificity of the enzyme.

The specificity of the interaction between PTP-PEST and p130 was addressed using the catalytic domain of PTP1B 20 (amino acid residues 1-321) in dephosphorylation reactions. When added at a similar molar concentration to that used for PTP-PEST, PTP1B was found to dephosphorylate fully and rapidly (within 15 minutes) most of the phosphotyrosinecontaining proteins present in the pervanadate-treated HeLa 25 In addition, the time course of dephosphorylation lysate. of p130 was not significantly more rapid than that of the other phosphotyrosine bands dephosphorylated by PTP1B. should be noted, however, that these in vitro dephosphorylation results are not truly illustrative of the substrate specificity of PTP1B in vivo for several reasons. First, only the isolated catalytic subunit was used in this particular experiment. Furthermore, in vivo substrate specificity may be quite different due to the intracellular distribution of both the PTP and potential substrates.

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is, in vitro dephosphorylation experiemnts may utilize substrates which the PTP is capable of dephosphorylating but which it would not have access to in vivo. The phenomenon of differing substrate specificity depending upon different physiologic contexts is illustrated by a comparison of this data with the in vivo PTP1B work described above, wherein PTP1B showed specificity for only three proteins.

Identification of Phosphotyrosine-Containing p130 Protein as p130 cas by Substrate Trapping

Pervanadate-treated HeLa cell lysate was fractionated by anion exchange chromatography and aliquots of the fractions were analyzed by SDS-PAGE followed by immunoblotting with anti-phosphotyrosine or anti-p130 cas antibodies. Aliquots of all samples analyzed were then 15 incubated with an affinity matrix containing a substrate trapping PTP-PEST mutant, comprising full length PTP-PEST in which Asp199 is changed to alanine (D199A), bound to covalently coupled protein A-Sepharose/antibody (AG25) beads. Proteins associated with PTP-PEST were then analyzed 20 by SDS-PAGE followed by immunoblotting with antiphosphotyrosine or anti-p130 cas antibodies.

Anti-phosphotyrosine immunoblotting of the column fractions showed that the pl30 phosphotyrosine band eluted as a single peak in fractions 11-14 (approx. 0.3 M NaCl). In view of the abundance of tyrosine phosphorylated pl30 in HeLa lysates, it appeared likely that p130 represents a previously identified phosphotyrosine-containing 130 kDa protein. Several potential candidates were identified in the literature, including the focal adhesion kinase pl25 FAK, ras-GAP, gp130 and pl30 cas. Of these candidates, p130 cas has been identified as a particularly prominent phosphotyrosine band in a wide variety of systems, including v-crk (Mayer and Hanafusa, Proc. Natl. Acad. Sci. USA 87: 2638-2642 (1990); Mayer et al., Nature 332:272-275 (1988))

and src (Kanner et al., Proc. Natl. Acad. Sci. USA 87:3328-3332 (1990); Reynolds et al., Mol. Cell. Biol. 9: 3951-3958 (1989)) transformed fibroblasts, integrin-mediated cell adhesion (Nojima et al., J. Biol. Chem. 270:15398-15402 (1995); Petch et al., J. Cell Science 108:1371-1379 (1995); Vuori and Ruoslahti, J. Biol. Chem. 270:22259-22262 (1995)) and PDGF stimulated 3T3 cells (Rankin and Rozengurt, J. Biol. Chem. 269:704-710 (1994)).

Therefore, the possibility that the p130

phosphotyrosine band corresponds to p130 was tested by immunoblotting the Mono Q fractions using an antibody to p130 cas. The 130 kDa band corresponding to p130 cas eluted in the same fractions as the p130 tyrosine phosphorylated band, and displayed a similar apparent molecular weight, suggesting that they might represent the same protein. Furthermore, p130 immunoprecipitated from these fractions was found to be phosphorylated on tyrosyl residues.

A mutant form of PTP-PEST (D199A) was generated by site-directed mutagenesis, and the mutant enzyme was 20 purified following expression using recombinant baculovirus. When assayed using tyrosine phosphorylated RCM-Lysozyme as substrate, the purified mutant enzyme exhibited a specific activity which was approximately 10,000 fold lower than that of the wild type enzyme (Garton and Tonks, unpublished 25 data). This purified protein was bound to an affinity matrix comprised of an anti-PTP-PEST monoclonal antibody (AG25) covalently coupled to Protein A-Sepharose beads, then incubated with each of the Mono Q fractions. After 45 minutes of incubation, proteins associating with the mutant 30 PTP-PEST were collected by centrifugation, the beads were washed, and SDS-PAGE sample buffer was added. Associated proteins were then analyzed by immunoblotting using the monoclonal anti-phosphotyrosine antibody G104.

The mutant PTP-PEST protein was found to associate with a single phosphotyrosine-containing protein, the molecular weight (130 kDa) and Mono Q elution position (fractions 11-14) of which coincided with those of $p130^{\text{Cas}}$ Immunoblotting of the PTP-PEST-associated proteins using the $p130^{\,\mathrm{cas}}$ antibody demonstrated that the 130 kDa tyrosine phosphorylated protein trapped by the mutant PTP-PEST is indeed p130 cas. These data further support the hypothesis that p130 cas is a potential physiologically relevant substrate for PTP-PEST.

Determination of Structural Features of PTP-PEST Involved in Specific Interaction with Tyrosine Phosphorylated p130 cas

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The interaction between pl30 cas and PTP-PEST was investigated further in substrate trapping experiments using various purified mutant forms of PTP-PEST to precipitate proteins from pervanadate-treated HeLa lysates. affinity matrices were incubated with pervanadate-treated HeLa cell lysate, and proteins associated with the beads were analyzed by SDS-PAGE followed by immunoblotting with anti-phosphotyrosine or anti-pl30 cas antibodies.

The wild type full-length phosphatase was found to be incapable of stable association with tyrosine phosphorylated p130 cas, whereas both the PTP-PEST (D199A) mutant protein and a mutant lacking the active site cysteine residue (C231S) specifically precipitated p130 cas from the lysate. The inability of the wild type phosphatase to precipitate tyrosine phosphorylated p130 cas presumably reflects the transient nature of the normal interaction between PTP-PEST and tyrosine phosphorylated p130 cas, which is likely to be 30 concluded as soon as p130 cas is dephosphorylated by PTP-PEST.

Since the C-terminal 500 amino acids of PTP-PEST contain several proline-rich regions which resemble src. homology-3 (SH3) domain binding sequences, it appeared

plausible that the specificity of the interaction between PTP-PEST and p130 cas might depend to some extent on association of these segments with the SH3 domain of The possible contribution of the C-terminal segment of PTP-PEST in the observed specific interaction of PTP-PEST with p130 was therefore addressed in further substrate trapping experiments using GST fusion proteins containing the catalytic domain of PTP-PEST alone, in both wild type and mutant (D199A) forms. The mutant catalytic domain of PTP-PEST fused to GST was found to precipitate the $pl30^{\text{Cas}}$ phosphotyrosine band specifically, whereas both the wild type fusion protein and GST alone failed to precipitate The specific interaction between PTP-PEST and pl30 cas observed in these experiments therefore appears to be an intrinsic property of the catalytic domain of PTP-15 PEST, emulating the observed preference of the active PTP-PEST catalytic domain for dephosphorylation of pl30 cas in <u>vitro</u>.

Specificity of Interaction Between Mutant PTP-PEST and Tyrosine Phosphorylated p130 Cas

In view of the relative abundance of tyrosine phosphorylated p130 cas in the pervanadate-treated HeLa cell lysate, the possibility that the observed selective binding of PTP-PEST inactive mutant proteins to p130 was substrate-directed (reflecting the abundance of this potential substrate relative to the other phosphotyrosine-containing proteins present in the lysate) rather than enzyme-directed (reflecting a genuine substrate preference of PTP-PEST) was considered; this possibility was addressed in two ways. First, inactive mutant forms of the catalytic domain of PTP1B were used to trap potential substrates for this enzyme from the pervanadate-treated HeLa lysates. Again it was found that the wild type phosphatase was incapable of stable interaction with any phosphotyrosine-

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containing protein, whereas mutant variants of the PTP1B phosphatase domain (comprising Cys or Asp mutations analogous to those described above for PTP-PEST) associated with many tyrosine phosphorylated proteins. This was especially apparent for the aspartic acid mutant of PTP1B (D181A), which appeared to precipitate essentially all phosphotyrosine-containing proteins from the lysate with similar efficacy. These data emphasize the specific nature of the interaction between PTP-PEST and p130 cas, which appears to be a property peculiar to the PTP-PEST catalytic domain, rather than a feature shared by all PTP catalytic domains.

The specificity of the interaction between PTP-PEST and p130 was addressed further following pervanadate
15 treatment of several different cell lines (Wi38, 293, COS, MCF10A, C2C12, MvLu), yielding a different array of tyrosine phosphorylated proteins in each case; the resultant lysates were analyzed by SDS-PAGE followed by anti-phosphotyrosine immunoblotting. Aliquots were incubated with PTP-PEST

20 (D199A) affinity matrix or control matrix, and tyrosine phosphorylated proteins associating with PTP-PEST were analyzed by SDS-PAGE and immunoblotting with anti-phosphotyrosine or anti-p130 antibodies as described above.

In each case, the D199A mutant PTP-PEST protein precipitated a single broad phosphotyrosine band with an apparent molecular weight between 120 and 150 kDa in different cell lines, whereas the affinity matrix alone failed to precipitate any phosphotyrosine-containing protein. Immunoblotting of the precipitates with a pl30 antibody revealed that the protein precipitated from all cell lysates corresponded to pl30 cas; the observed molecular weight variation between different cell lines presumably reflects either species differences in the

alternatively spliced forms (Sakai et al., EMBO J. 13:3748-3756 (1994)).

The relative abundance of tyrosine phosphorylated p130 cas in the PTP-PEST precipitates appeared to correlate approximately with the abundance of p130 cas protein in the lysates (data not shown). Surprisingly, regardless of the abundance of tyrosine phosphorylated p130 cas in the lysates, p130 cas was invariably the only phosphotyrosinecontaining protein in the precipitates, even in 293 cell lysates which contained very little p130 cas protein but which displayed a wide variety of other abundantly tyrosine phosphorylated proteins. Similarly, when lysates of pervanadate-treated 293 cells (containing tyrosine phosphorylated p130 cas in amounts which are undetectable by anti-phosphotyrosine immunoblotting of the lysate) were 15 incubated with active PTP-PEST, no visible dephosphorylation of any phosphotyrosine band occurred (Garton and Tonks, unpublished data). These results indicate that the affinity of PTP-PEST for p130 cas is substantially greater than for any other substrate present, and further emphasizes the remarkable substrate selectivity of PTP-PEST for pl30 cas.

<u>Vanadate Inhibition of Tyrosine Phosphorylated p130</u> Cas <u>Association with Mutant PTP-PEST</u>

A consistent observation of this work was that, in

contrast to the inactive mutant PTP-PEST, the wild type
enzyme failed to associate in a stable complex with tyrosine
phosphorylated p130 cas, suggesting that the observed
association is active site-directed. In order to
investigate this possibility, mutant PTP-PEST (D199A) was

incubated with the PTP inhibitor vanadate at various
concentrations prior to addition of pervanadate-treated HeLa
cell lysate. The extent of association of p130 cas with
PTP-PEST was then analyzed. PTP-PEST affinity matrix,
comprising full length PTP-PEST (D199A) bound to covalently

coupled protein A-Sepharose/antibody (AG25) beads, was incubated for 10 minutes on ice in the presence of varying concentrations of sodium orthovanadate. The samples were then incubated with aliquots of pervanadate-treated HeLa cell lysate; associated proteins were analyzed by SDS-PAGE and immunoblotting with anti-phosphotyrosine or anti-pl30 antibodies. The activity of wild type PTP-PEST was also determined under the same conditions, using tyrosine phosphorylated ³²P-labelled RCM-lysozyme as substrate.

The association was found to be potently disrupted by 10 vanadate, with a concentration-dependence similar to that of vanadate inhibition of wild type PTP-PEST, and complete disruption being observed at 10 mM vanadate. Since PTP inhibition by vanadate presumably results from a direct interaction of vanadate with the active site cysteine 15 residue of the enzyme (Denu et al., Proc. Natl. Acad. Sci. USA 93:2493-2498 (1996)), this result supports the hypothesis that the stable association of mutant PTP-PEST with tyrosine phosphorylated p130 cas is mediated by direct interactions between active site residues within PTP-PEST, 20 in particular the active site cysteine residue, and phosphotyrosine moieties within p130 cas.

Association of Endogenous p130 cas with Transfected Mutant PTP-PEST in COS Cells

The work described above strongly suggests that

p130 cas represents a potential physiologically significant substrate for PTP-PEST. In order to assess whether PTP-PEST interacts with p130 in intact cells, COS cells were transfected with plasmids encoding wild type or mutant forms of PTP-PEST (D199A or C215S). The cells were treated with pervanadate 30 minutes prior to lysis, PTP-PEST proteins were immunoprecipitated, and associated tyrosine phosphorylated proteins were analyzed by antiphosphotyrosine immunoblotting of the resultant

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precipitates. Lysates were also incubated with covalently coupled protein A-Sepharose/anti-PTP-PEST (AG25) beads and associated proteins were analyzed by SDS-PAGE and immunoblotting with anti-phosphotyrosine antibody.

Under these conditions, the phosphotyrosine-containing band corresponding to p130 was again unique in its ability to associate with the C231S PTP-PEST protein, indicating that p130 can be specifically selected by PTP-PEST as a substrate in an intracellular context in the presence of a large number of alternative possible substrates. Neither the wild type nor the D199A form of PTP-PEST was capable of a stable interaction with tyrosine phosphorylated p130 cas in pervanadate-treated COS cells.

The binding of both wild type and D199A PTP-PEST to tyrosine phosphorylated p130 cas under these conditions is 15 most likely prohibited by the presence of pervanadate bound to the active site cysteine residue of PTP-PEST (Denu et al., Proc. Natl. Acad. Sci. USA 93:2493-2498 (1996)), which effectively excludes the binding of phosphotyrosine residues of p130 Cas The ability of the C231S mutant PTP-PEST to 20 associate in a stable complex with p130 cas in the presence of pervanadate suggests that this mutant protein is largely unaffected by pervanadate, indicating that the normal mode of inhibition of PTPs by vanadate ions depends critically on direct interactions between vanadate and the thiolate anion 25 of the PTP active-site cysteine residue. These observations therefore lend further support to the existence of an exclusive interaction between PTP-PEST and pl30 cas which appears to be entirely active site-directed, and therefore reflects the genuine, inherent, highly restricted substrate preference of PTP-PEST for pl30 cas

Results described herein implicate p130^{cas} as a physiologically relevant substrate for PTP-PEST.
Furthermore, the observed stringency and exclusivity of the interaction between PTP-PEST and p130^{cas} in a wide variety

of cell lines suggest that p130 cas may be a unique high affinity substrate for PTP-PEST, although the possibility that other significant PTP-PEST substrates may exist cannot be excluded at present. In particular, it is unclear 5 whether pervanadate-treated cells display a complete spectrum of all possible tyrosine phosphorylated proteins; in fact this appears unlikely since pervanadate treatment presumably results only in an increase in tyrosine phosphorylation of proteins which are to some extent constitutively phosphorylated, but which are normally 10 rapidly dephosphorylated, within the cell. Potential substrates lacking from pervanadate-treated cells therefore presumably include substrates of protein tyrosine kinases (PTKs) which are normally present in an inactive state, such as ligand-stimulated receptor PTKs, and the recently described calcium regulated kinase PYK2 (Lev et al., Nature 376: 737-745 (1995)). Regardless of these considerations, the ability of PTP-PEST to select p130 cas exclusively as a substrate from lysates of several different cell lines, containing a combined total of at least one hundred 20 different potential substrates (many of which presumably contain multiple sites of phosphorylation), clearly demonstrates that the substrate specificity of PTP-PEST is highly restricted.

Many intracellular PTPs are limited in their substrate availability due to strict confinement within a particular subcellular location; examples include PTP1B, which is localized to the cytoplasmic face of the endoplasmic reticulum (Frangioni et al., Cell 68:545-560 (1992)), and 30 TCPTP which is either nuclear (Tillmann et al., Mol. Cell. Biol. 14:3030-3040 (1994)) or localized to the endoplasmic reticulum, depending upon which alternative spliced form is expressed (Lorenzen et al., J. Cell Biol. 131:631-643 Alternatively, certain PTPs appear to be highly regulated, requiring activation before appreciable activity WO 98/04712 PCT/US97/13016

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can be demonstrated. For example, the SH2 domain-containing PTPs, SHP1 and SHP2, display relatively low activity in vitro, but can be considerably activated by several mechanisms including C-terminal truncation (Zhao et al., J. Biol. Chem. 268:2816-2820 (1993)), addition of certain phospholipids (Zhao et al., Proc. Natl. Acad. Sci. USA 90:4251-4255 (1993)), or SH2 domain-mediated binding of appropriate phosphotyrosine-containing peptides (Lechleider et al., J. Biol. Chem. 268:21478-21481 (1993)).

However, PTP-PEST exhibits high specific activity in 10 vitro (35,000 U/mg), and is a predominantly (90-95%) soluble PTP within cells (Garton and Tonks, unpublished data); in principle, therefore, it may act potently on any substrate accessible to the cytoplasm. This accessibility may partly 15 underlie the necessity for PTP-PEST to possess an inherently constrained substrate specificity. The demonstration that mutant PTP-PEST is capable of exclusively associating with $p130^{\text{Cas}}$ in an intracellular context in the presence of many other tyrosine phosphorylated proteins, is an indication that the narrow substrate specificity of the enzyme may result in PTP-PEST having a negligible influence on the phosphorylation state of the majority of tyrosine phosphorylated proteins within the cell, even though those substrates are largely accessible to PTP-PEST.

The role of p130 cas in cellular transformation by the v-crk and v-src oncogenes is unclear, although there is a general correlation between the level of tyrosine phosphorylation of p130 and the degree of transformation in cells expressing different forms of crk or src (Kanner et al., EMBO J. 10:1689-1698 (1991); Mayer and Hanafusa, J. Virol. 64:3581-3589 (1990)). Furthermore, enhanced tyrosine phosphorylation of p130 has also been observed in cells transformed by c-Ha-ras and by ornithine decarboxylase overexpression (Auvinen et al., Mol. Cell. Biol. 15:6513-6525 (1995)). Expression of antisense cDNA encoding

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p130^{Cas} in these cells results in a partial reversion of the transformed phenotype. These observations suggest that aberrant tyrosine phosphorylation of p130^{Cas} is a common feature of cells transformed by several disparate mechanisms and that p130^{Cas} may be required for full manifestation of the transformed state. Dephosphorylation of p130^{Cas} by PTP-PEST is therefore a potentially important regulatory mechanism for counteracting the transforming effects of various oncogenes.

Tyrosine phosphorylation of p130 cas has been observed 10 in fibroblasts following integrin-mediated cell adhesion to extracellular matrix proteins (Nojima et al., J. Biol. Chem. 270:15398-15402 (1995); Petch et al., J. Cell Science 108:1371-1379 (1995); Vuori and Ruoslahti, J. Biol. Chem. 270:22259-22262 (1995)). Under these conditions, using an 15 antibody (4F4) that predominantly recognizes tyrosine phosphorylated p130 cas (Kanner et al., EMBO J. 10:1689-1698 (1991); Petch et al., J. Cell Science 108:1371-1379 (1995)) it was shown that phosphorylated p130 cas is localized to focal adhesions (Petch et al., J. Cell Science 108:1371-1379 (1995)), whereas fractionation studies have demonstrated that the normal cellular location of the majority of nonphosphorylated p130 cas is the cytosol (Sakai et al., EMBO <u>J. 13</u>:3748-3756 (1994)). Furthermore, in crk-transformed fibroblasts, tyrosine phosphorylated p130 cas is detected 25 only in insoluble fractions (Sakai et al., EMBO J. 13:3748-3756 (1994)), suggesting that both cell adhesion- and transformation-mediated phosphorylation of p130 cas is associated with redistribution of the protein from the cytosol to focal adhesions.

It is plausible that the redistribution of tyrosine phosphorylated p130 may be driven by its association with FAK, which is constitutively associated with focal adhesions due to its C-terminal focal adhesion targeting domain (Hildebrand et al., J. Cell Biol. 123:993-1005

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(1993); Schaller et al., Proc. Natl. Acad. Sci. USA 89:5192-5196 (1992)). The sequestration of tyrosine phosphorylated p130 cas in focal adhesions both in transformed cells, and following integrin-mediated cell adhesion, strongly suggests a role for p130 cas in signalling events in this region of the cell. One consequence of the redistribution of tyrosine phosphorylated p130 cas is likely to be that, in addition to localizing p130 cas to a region of the cell containing abundant protein tyrosine kinase activity, the phosphorylated protein will be relatively inaccessible to the cytosolic phosphatase PTP-PEST. This raises the possibility that the role of PTP-PEST in dephosphorylating pl30 cas may be to prevent inappropriate tyrosine phosphorylation of the cytosolic pool of p130 cas, thus preventing formation of signalling complexes assembled around tyrosine phosphorylated pl30 cas in inappropriate cellular locations.

Several mitogenic factors potently stimulate tyrosine phosphorylation of pl30 cas. These include agents acting through heterotrimeric G protein-coupled receptors such as 20 lysophosphatidic acid (Seufferlein and Rozengurt, J. Biol. Chem. 269:9345-9351 (1994)), bradykinin (Leeb-Lundberg et al., J. Biol. Chem. 269: 24328-24344 (1994)), and bombesin (Zachary et al., J. Biol. Chem. 267:19031-19034 (1992)), as well as growth factors that activate receptor tyrosine 25 kinases, namely PDGF (Rankin and Rozengurt, J. Biol. Chem. 269:704-710 (1994)), EGF and NGF (Ribon and Saltiel, J. Biol. Chem. 271:7375-7380 (1996)). These observations suggest roles for p130 cas in regulation of mitogenic signalling pathways, presumably involving assembly of 30 signalling complexes based on tyrosine phosphorylated pl30 cas. The identities of the proteins involved in these complexes are not established, but are likely to include SH2 domain-containing adaptor proteins such as crk (Ribon and Saltiel, J. Biol. Chem. 271:7375-7380 (1996)), and its 35

associated proteins (Feller et al., Oncogene 10:1465-1473 (1995); Hasegawa et al., Mol. Cell. Biol. 16:1770-1776 (1996); Knudsen et al., J. Biol. Chem. 269:32781-32787 (1994); Matsuda et al., Mol. Cell. Biol. 14: 5495-5500 (1994); Tanaka et al., Proc. Natl. Acad. Sci. USA 91:3443-3447 (1994)). Therefore tyrosine phosphorylation and dephosphorylation of p130 potentially plays a central role in regulating the formation of such complexes, thereby influencing downstream events in mitogenic signalling.

10 <u>Equivalents</u>

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Those skilled in the art will recognize, or be able to ascertain, using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following claims:

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CLAIMS

We claim:

- 1. A protein tyrosine phosphatase wherein the invariant aspartate residue is replaced with an amino acid (e.g. alanine) which:
 - a) does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute; or
- b) is selected from the group consisting of:
 alanine, valine, leucine, isoleucine, proline,
 phenylalanine, tryptophan, methionine, glycine,
 serine, threonine, cysteine, tyrosine, glutamine,
 lysine, arginine and histidine, and wherein the
 phosphatase binds to a tyrosine phosphorylated
 substrate and is catalytically attenuated.
- 2. A protein tyrosine phosphatase according to claim 1(a) which is selected from the group consisting of: PTP1B (and wherein for example the invariant aspartate residue is located at position 181), PTP-PEST (and wherein for example the invariant aspartate residue is located at position 199), PTPτ, MKP-1, DEP-1, PTPμ, PTPX1, PTPX10 and PTPH1.
- A protein tyrosine phosphatase according to claim 1(b) which is a PTP-PEST phosphatase in which the amino
 acid at position 231 is replaced with a serine residue.
 - 4. A method of identifying a tyrosine phosphorylated protein which is a substrate of a protein tyrosine phosphatase, comprising the steps of:
- a) combining at least one tyrosine phosphorylated protein with at least one protein tyrosine

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b)

phosphatase in which the invariant aspartate residue is replaced with an amino acid (e.g. alanine) which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, under conditions appropriate for formation of a complex between the tyrosine phosphorylated protein and the protein tyrosine phosphatase, thereby producing a combination; and determining the presence of absence of a complex in the combination wherein the presence of a complex in the combination indicates that the tyrosine phosphorylated protein is a substrate of the protein tyrosine phosphatase with which it forms a complex.

- A method according to claim 4, wherein the protein tyrosine phosphatase is as defined in any one of claims 1-3.
- A method according to claim 4 or claim 5 wherein the 6. tyrosine phosphorylated protein is selected from the 20 group consisting of: pl30^{cas}, the EGF receptor, p210 bcr:abl, MAP kinase and the insulin receptor.
- A kit for identifying a tyrosine phosphorylated 7. protein substrate of a protein tyrosine phosphatase comprising: 25
 - at least one protein tyrosine phosphatase in a) which the invariant aspartate residue is replaced with an amino acid (e.g. alanine) which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute; and

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- b) ancillary reagents suitable for use in detecting the presence or absence of a complex between the protein tyrosine phosphatase and a tyrosine phosphorylated protein,
- wherein for example the protein tyrosine phosphatase is as defined in any one of claims 1-3.
 - 8. A method of identifying an agent which alters the interaction between a protein tyrosine phosphatase and a tyrosine phosphatase, comprising the steps of:
 - a) identifying a tyrosine phosphorylated protein which is a substrate of a protein tyrosine phosphatase;
 - b) combining the tyrosine phosphorylated protein and the protein tyrosine phosphatase and an agent to be tested under conditions suitable for interaction between the tyrosine phosphorylated protein and the protein tyrosine phosphatase, thereby forming a combination;
 - c) determining the amount of enzymatic activity in the combination; and
 - d) comparing the amount of enzymatic activity determined in (c) with the amount of enzymatic activity in the absence of the agent to be tested, under conditions suitable for interaction between the tyrosine phosphorylated protein and the protein tyrosine phosphatase,

wherein a difference in the enzymatic activity indicates that the agent alters the interaction between the protein tyrosine phosphatase and the tyrosine phosphorylated protein.

9. A method of identifying an agent which alters the interaction between a protein tyrosine phosphatase and a tyrosine phosphorylated protein which is a substrate

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of the protein tyrosine phosphatase, comprising the steps of:

- a) identifying a tyrosine phosphorylated protein which is a substrate of a protein tyrosine phosphatase;
- b) combining the tyrosine phosphorylated protein, a protein tyrosine phosphatase in which the invariant aspartate residue is replaced with an amino acid (e.g. alanine) which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, and an agent to be tested, under conditions suitable for interaction between the tyrosine phosphorylated protein and the protein tyrosine phosphatase, thereby forming a combination;
- c) determining the extent of binding between the tyrosine phosphorylated protein and the protein tyrosine phosphatase in the combination; and
- d) comparing the extent of binding determined in (c) with the extent of binding in the absence of the agent to be tested, under conditions suitable for interaction between the tyrosine phosphorylated protein and the protein tyrosine phosphatase,
- wherein a difference in the extent of binding indicates that the agent alters the interaction between the protein tyrosine phosphatase and the tyrosine phosphatase and the tyrosine phosphorylated protein.
- 30 10. A method according to claim 8 or claim 9 wherein if the amount of enzymatic activity or the extent of binding, respectively, is:
 - a) greater in the presence of the agent to be tested than in the absence of the agent, then the agent

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enhances the interaction between the protein tyrosine phosphatase and the tyrosine phosphorylated protein; or

- b) less in the presence of the agent to be tested than in the absence of the agent, then the agent inhibits the interaction between the protein tyrosine phosphatase and the tyrosine phosphorylated protein.
- 11. A protein tyrosine phosphatase (e.g. as defined in any one of claims 1-3) for use in therapy, prophylaxis or diagnosis, for example in:
 - a) the treatment of conditions in which a reduction in the activity of a tyrosine phosphorylated protein is indicted; and/or
- 15 b) reducing the activity of a tyrosine phosphorylated protein; and/or
 - c) a method of reducing the activity of a tyrosine phosphorylated protein, comprising administering to a mammal (e.g. a human) a protein tyrosine phosphatase in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, whereby formation of the complex reduces the activity of the tyrosine phosphorylated protein; and/or
 - d) the treatment of conditions in which a reduction in the transforming effects of oncogenes associated with p130^{cas} phosphorylation is indicated; and/or
 - e) reducing the transforming effects of oncogenes associated with p130^{cas} phosphorylation; and/or
 - f) a method of reducing the transforming effects of oncogenes associated with pl30^{cas} phosphorylation

		comprising administering to a mammal (e.g. a
:		human) a protein tyrosine phosphatase which is
		PTP-PEST or PTP-PEST in which the invariant
		aspartate residue is replaced with an amino acid
		which does not cause significant alteration of
5		the Km of the enzyme but which results in a
		reduction in Kcat to less than 1 per minute,
		whereby the PTP binds to and/or dephosphorylates
		p130 ^{cas} and reducing the transforming effects of
10		oncogenes associated with p130 cas phosphorylation;
•		and/or
	g)	the treatment, therapy, diagnosis or prophylaxis
		of cancer (for example cancers associated with
		p130 ^{cas} phosphorylation); and/or
15	h)	the treatment of conditions associated with
		oncogneic activity (e.g. with v-crk, v-src and/or
		c-Ha-ras activity); and/or
	įi)	the treatment of conditions in which a reduction
	:	in the formation of signalling complexes
20		associated with p130 ^{cas} is indicted; and/or
	j)	reducing the formation of signalling complexes
•		associated with pl30 ^{cas} ; and/or
	k)	a method of reducing the formation of signalling
	•	complexes associated with pl30cas comprising
. 25	•	administering to a mammal (e.g. a human) a
	•	protein tyrosine phosphatase which is PTP-PEST or
		PTP-PEST in which the invariant aspartate residue
		is replaced with an amino acid which does not
	•	cause significant alteration of the Km of the
30		enzyme but which results in a reduction in Kcat
50	:	to less than 1 per minute, whereby the PTP binds
		to and/or dephosphorylates p130 ^{cas} and reducing the
· · · · · · · · · · · · · · · · · · ·		formation of signalling complexes associated with
	•	p130 ^{cas} , and/or
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q)

- the prevention of the induction of mitogenic pathways;
- m) the treatment of conditions in which the prevention of the induction of mitogenic pathways is indicated;
- n) the treatment of conditions in which a reduction in the cytotoxic effects associated with protein tyrosine phosphatase administration or overexpression is indicated; and/or
- o) reducing the cytotoxic effects associated with protein tyrosine phosphatase administration or over-expression; and/or
 - p) a method for reducing the cytotoxic effects associated with protein tyrosine phosphatase administration or over-expression comprising administering to a mammal (e.g. a human) a protein tyrosine phosphatase in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, in place of a corresponding wild type protein tyrosine phosphatase; and/or

in a method according to any one of claims 13-20.

- 25 12. Use of a protein tyrosine phosphatase for the manufacture of a medicament for treatment, prophylaxis or diagnosis (for example for use in the treatments defined in claim 11).
- 13. A method of reducing the activity of a tyrosine
 phosphorylated protein, comprising administering to a
 mammal (e.g. a human) a protein tyrosine phosphatase
 in which the invariant aspartate residue is replaced
 with an amino acid which does not cause significant

alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, whereby formation of the complex reduces the activity of the tyrosine phosphorylated protein.

- 5 14. A method according to claim 13, wherein the tyrosine phosphorylated protein is selected from the group consisting of: p130^{cas}, the EGF receptor, p210 bcr:abl, MAP kinase and the insulin receptor.
- 15. a method according to claim 13, wherein the protein tyrosine phosphatase is selected from the group consisting of: PTP1B, PTP-PEST, PTPτ, MKP-1, DEP-1, PTPτ, PTPX1, PTPX10 AND PTPH1.
- A method of reducing the transforming effects of oncogenes associated with p130cas phosphorylation comprising of administering to a mammal (e.g. a human) 15 a protein tyrosine phosphatase which is PTP-PEST or PTP-PEST in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 20 per minute, whereby the PTP binds to and/or dephosphorylates p 130cas, thereby negatively regulating the downstream effects of pl30^{cas} and reducing the transforming effects of oncogenes associated with p130^{cas} phosphorylation. 25
 - 17. A method according to claim 16, wherein the oncogene is selected from the group consisting of: v-crk, v-src and c-Ha-ras.

- 18. A method of reducing formation of signalling complexes associated with pl30^{cas} comprising administering to a mammal (e.g. a human) a protein tyrosine phosphatase which is PTP-PEST OR PTP-PEST in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, whereby the PTP binds to and/or dephosphorylates pl30^{cas}, thereby negatively regulating the downstream effects of pl30^{cas} and reducing the formation of signalling complexes associated with pl30^{cas}.
 - 19. A method according to claim 18, which prevents the induction of mitogenic pathways.
- 15 20. A method of reducing cytotoxic effects associated with protein tyrosine phosphatase administration or over expression, comprising administering to a mammal (e.g. a human) a protein tyrosine phosphatase in which the invariant aspartate residue is replaced with an amino acid which does not cause significant alteration of the Km of the enzyme but which results in a reduction in Kcat to less than 1 per minute, in place of a corresponding wild type protein tyrosine phosphatase.

QKSRGVVMLNRVM QKTKAVVMLNRIV	HNVEVIVATIONLY	ONTGILVMITNLV	QRVALIVMITNLV	QRIATVVMMTRLE	ENTASIIMVTNLV	QNTATIVMVINLK	OKSALLVMLINLK	QKAIVIVMTKEV FNSPVIVMTFKEV	ENSRVIVMTTREV	QNVHNIVMVIQĆV	SNSRAIVMLTRCF	QOSHILVMLTNCM	ERTPITAMITNIE	OHLEITVILTNLE	OKVLVIVMITRFE	YNVVIIVMACKEF	OKLISTIVMUTTLT	NRCRIIVMLSRES	VENIGITAMESSLF	SGINGVIVEDINE	THINGE A VALUATION	HIND LIT VILLE NEW RINGS IN THE PART TO COME WINDS	WKSHTT/MTTEVO	YHCTSVVMLNDVD	RKVKVI WLTELV	QSVTTLVMISEIG	HNAQLVWIPDGQ	HNAQII VALPDNQ GIYYYA LESI	CHARGE V VICTORIA	1		ta Sc	•
YILTGGPLPNTCCH <i>FW</i> EMVWD	YIAAGGPLKSTAEDFWRMIWE	TATAGETIKSTFED FWRMINE	FIGTGEPLPDTFDCFWRMIWE	YIATGGPLPETMGD <i>FW</i> RMVWE	YIATGGPMQETIYD FWRMVWH	FIAAGGPKEETVNDFWRMIWE	FIAAGGPKOETVNDFWRMVWE	TATAAGGPRDETVUDFWRMIWE	YTASOGCI, BATVND FROMAWO	TIVIGELPGIKDDFWKMVWE	RIVIGEPLHSTRDD FWRMCWE	SFIATGGPLPQTVGDFWRLVWE	/YIATGGPIVSTVVDFWRMVWQ	CFICAGGPMESTIDD FWRMIWE	AYIGTGGPLENTYRDFWLMVWE	AYVATGGPLANTVIDFWRMVWE	KYIATGGPLPHTCAOFWOVVWD	DFICT GEPLPNTIAD FWRWWE	NYIACGGSISRSISDFWHMVWDN	KYIACGAPKPGTLLDFWEMVWHN	AY IAAGGPLKSTAEDFWKMIWE	AY IATGEFLAESTEDFWKMLWE	SILASQGFULDILLEDFWilling	AFIVICHPLPNIVKDFWRLVLD	MMIAAGGPLKETIGDFWOMIFQ	TFIIAQDPFENTIGDFWRMISE	EFIITGH PLLHTIKDFWRMIWD	EFIITGH PLPHTTKDFWRMIWD	DETVICE FAMALLIANT WORK TO	RT. LACGY PLOSOLESHERMLAE	O.S.	p2 + c2	
KMEE	DGYN	NISON	DGXO	DGYR	HXDC	NGYO	DGYK		. •	PGNN	NHDd		RGYNG	IGYK	DGYK	KGVY	PAAN L	DGA			DGXN	DGYR	DGYK	DSYK	SKYINASFV MSYW	TYINASFI EGYD	MGYY	MGYY	HGFK K			pl Figure 1A	
PTP1B	PTP_xi_D1 DGKLT	PTP_zela_D1 D	PIF_Gamma_D DSMS PTP99A D1 KKNL	PGS	PTP mu D1 TNS	PTP_alpha_D1 PDS	PTP_opsilon_D PCS	. AGS	SH. PTP2 EPV	יי ה ה ה ה	EGS BGS	SAP.1	PTP STEP DPL S	D1 QTT	TQT		Hum_PTPH1 INA	PTP1	t_pyp1			,	_D2	2n_02	SEET	2 ENS	2	_D2	7	Yarsinia_PTP	PTP1Bseq.no.	PTP1B66	
	PTP1B DYINASLI KMEBAQRSYILIGGPLPNTCGHFWEMVWE DIBEAQRSYILIGGPLPNTCCHFWLMVWQ	TCPTP TCPTP TCPTP DGKLTDYINANYV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RPKAYIAAGGPLKSTAEDFWRMIWE	FTP1B DyINASLI DyINASLI DIEEAQRSYILTGGPLPNTCGHFWEMVWE DYINASLV DIEEAQRSYILTGGPLPNTCCHFWLMVWQ DYINASLV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIATGGPLKSTEEDFWRMIWE DGYN KAKAYIATGGPLKSTEEDFWRMIWE	PTP1B DyINASLI DyINASLI DYINASLY DGEAQRSYILITGGPLPNTCCHFWLMVWE DYINASLY DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIAAGGPLKSTFEDFWRMIWE DGYN KAKAYIATGGPLKSTFEDFWRMIWE DGYN KAKAYIATGGPLKSTFEDFWRMIWE DGYN KAKAYIATGGPLKSTFEDFWRMIWE DTP99A D1 KKNL DYINANFI DGYQ KGHAFIGTGGPLPDTFDCFWRMIWE	TCPTP TCPTP DyINASLI DyINASLI DGKLTDYINANIV PTP_zela_D1 DGKLTDYINANIV PTP_gamma_D DSKHSDYINANIV PTP_gamma_D KKNL DYINANIY DGYN KAKAYIATGGPLFURNIME DGYN KAKAYIATGGPLKSTAEDFWRMIWE DGYN KAKAYIATGGPLKSTFEDFWRMIWE DGYN KAKAYIATGGPLFUFFRENIME DGYN KAKAYIATGGPLFDTFDCFWRMIWE DGYN KGHAFIGTGGPLPDTFDCFWRMIWE DGYN KGHAFIGTGGPLPDTFDCFWRMIWE DGYN KGHAFIGTGGPLPDTFDCFWRMIWE DGYN KONAXIATGGPLPDTFDCFWRMIWE DGYN KONAXIATGGPLPDTFDCFWRMIWE DGYN KONAXIATGGPLPDTFDCFWRWIWE	TCPTP PTP1B DyIMASLI DyIMASLI DYIMASLY DGKLTDYINANYV PTP_zela_D1 DGKLTDYINANYV PTP_gamma_D PTP_gamma_D PTP_gamma_D ETPP99A_D1 KKNL DYIMANYV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIATGGPLKSTFEDFWRMIWE DGYN KAKAYIATGGPLKSTFEDFWRMIWE DGYN KAKAYIATGGPLPDTFDCFWRMIWE DGYN KAKAYIATGGPLPDTFDCFWRMIWE DGYN KAKAYIATGGPLPDTFDCFWRMIWE DGYN KONAYIATGGPLPTFDCFWRMIWE DGYN KONAYIATGGPLPTFDCFWRWIWE DGYN RONAYIATGGPLPTTYDFWRWWH TNS DYINGNYI DGYN RONAYIATGGPLPTTYDFWRWWH	PTP1B DyINASLI DYINASLI DIEEAQRSYILITGGPLPNTCGHFWEMVWE DIEEAQRSYILITGGPLPNTCCHFWLMVWQ PTP_zela_D1 DGKLTDyINANYV PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_GAD1 PGS DYINANYV DGYN RAKAYIATGGPLKSTAEDFWRMIWE DGYN RAKAYIATGGPLFBDFWRMIWE DGYN RAKAYIATGGPLPDTFDCFWRMIWE DGYN RONAYIATGGPLPDTFDCFWRMIWE DGYN RONAYIATGGPLPTFMCFWRWYWE DGYN RONAYIATGGPLPTFWRWWWH DGYN RONAYIATGGPLPTFWRWWH RONAYIATGGPLPTFWRWWH RONAYIATGGPRETTWDFWRWWH RONAYIATGGPRETTWDFWRWH RONAYIATGGPRETTWDFWRWWH RONAYIATGGPRETTWDFWRWH RONAYIATGGPRETTWDFWRWWH RONAYIATGGPRETTWDFWRWWH RONAYIATGGPRETTWDFWRWH RONAYIATGGRETTWDFWRWH RONAYIATGGPRETTWDFWRWH	PTP1B DyINASLi DyINASLi DGKLTDYINANIV PTP_zela_D1 DGKLTDYINANIV PTP_gamma_D PTP_gamma_D DSKHSDYINANIV PTP_gamma_D DSKHSDYINANIV PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PGYN RPKAYIAAGGPLFWERMIWE DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RKNIL DYINANIV DGYN ROYN ROYN RENNKFIAAGGPKEETVNDFWRMIWE DGYK BKNKFIAAGGPKEETVNDFWRMIWE DGYK BKNKFIAAGGPKETVNDFWRMIWE DGYK BKNKFIAAGGPKGETVNDFWRMIWE DGYK BKNKFIAAGGPKGETVNDFWRMIWE DGYK BKNKFIAAGGPKGETVNDFWRMIWE DGYK BKNKFIAAGGPKGETVNDFWRMIWE DGYK BKNKFIAAAGGPKGETVNDFWRMIWE DGYK BKNKFIAAAGGPKGTANDFWRMIWE DGYN BKNKFIAAAGGPKGTANDFWRMIWE DGYK BKNKFIAAAGGPKGTANDFWRMIWE DGYK BKNKFIAAAGGPKGTANDFWRMIWE DGYK BKNKFIAAAGGPKGTANDFWRMIWE DGYK BKNKFIAAAGGPKGTANDFWRMIWE DGYK BKNKFIAAAGGPKANIWE DGYK BKNKFIAAAGGPKANIWE DGYN BKNKFIAAAGGPKANIWE DGYN BKNKFIAAAGGPKANIWE BKNKFIAAAAGGPKANIWE BKNKFIAAAGGPKANIWE BKNKFIAAAGGPKANIWE BKNKFIAAAGGPK	PTP1B DyINASLI DyINASLI DYINASLY DGYN RPKAYIAAGGPLPNTCCHFWLMVWQ PTP_zela_D1 DGKLTDYINANYV PTP_gamma_D PGKLTDYINANYV PTP_gamma_D PGS DYINANYV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIATGGPLPDTFDCFWRMIWE DGYN KAKAYIATGGPLPDTFDCFWRMIWE DGYN KAKAYIATGGPLPDTFDCFWRMIWE DGYN KAKAYIATGGPLPDTFDCFWRMIWE DGYN KQNAYIATGGPLPETMGDFWRWIWE DGYN KQNAYIATGGPLPETMGDFWRWIWE DGYN RQNAYIATGGPLPETMGDFWRWIWE PTP_alpha_D1 PDS DYINANYI DGYR KQNAYIATGGPLPETMGDFWRWIWE DGYR KQNAYIATGGPLPETMGDFWRWIWE DGYR KCHKFIAAGGPKGETVNDFWRWIWE DGYR EKNKFIAAGGPKGETVNDFWRWIWE DGYK EKNKFIAAGGPROETVDDFWRMIWE DGYK ERNKFIAAGGPROETVDDFWRMIWE DGYK ERNKFIAAGGPROETVDFWRMIWE DGYK ENDELEDFROETVDFWRMIWE DGYK ENDELEDFROETVDFWRMIWE DGYK ENDELEDFROETVD	PTP1B DyINASLI DyINASLI DYINASLI DGEAQRSYILITGGPLPNTCGHFWEMVWE DIEEAQRSYILITGGPLPNTCCHFWLMVWQ PTP_zela_D1 DGKLTDYINANYV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIAAGGPLKSTAEDFWRMIWE DGYN RAKAYIAATGGPLKSTAEDFWRMIWE DGYN RAKAYIAATGGPLEATVNDFWRMIWE DGYN RAKAYIAATGGPLATVNDFWRMIWE DGYN RAKAYIAATGG	PTP1B DyINASLI DyINASLI DGKLTDJINANY DGXN RPKAYIAAGGPLKSTAEDFWRMIWE PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PGS DYINANY PTP_gamma_D FKNL DYINANY PGS DYINANY PTP_alpha_D1 PGS DYINANY PTP_alpha_D1 PGS DYINASYI PGS DYINASYI DGYR KQNAYIATGGPLPDTFDCFWRMIWE DGYR KQNAYIATGGPLPDTFDCFWRMIWE DGYR KONAYIATGGPLPDTFDCFWRMIWE DGYR KQNAYIATGGPLPTFDCFWRMIWE DGYR KQNAYIATGGPLPTFNCFWRWIWE DGYR KQNAYIATGGPLPTFNCFWRWIWE DGYR KONAYIATGGPLPTFNCFWRWIWE DGYR KONAYIATGGPLPTFNCFWRWIWE DGYR KONAYIATGGPLPTFNCFWRWIWE DGYR KONAYIATGGPLPTFNCFWRWIWE DGYR EKNKFIAAGGPRETVNDFWRWIWE DGYR ERRKYIAAGGPRETVNDFWRWIWE SH. PTP2 EPV SDYINANYIKNOLL GPDE NAKTYIASGGCLEATVNDFWRWIWE PCS DYINANYIKNOLL GPDE NAKTYIASGGCLEATVNDFWRWIWE PCS DYINASYI	PTP1B DyINASLI DyINASLI DJINASLI DJINASLI DJINASLI DJINASLI DJINASLI DJINASLI DJINASLI DGKLITDYINANY DGYN RPKAYIAAGGPLKSTAEDFWRMIWE DGYLTDYINANY DGYN RAKAYIATGGPLKSTREDFWRMIWE DGYN RAKAYIATGGPLKSTFEDFWRMIWE DGYN RAKAYIATGGPLKSTFEDFWRMIWE DGYN RAKAYIATGGPLKSTFEDFWRMIWE DGYN RAKAYIATGGPLKSTFEDFWRMIWE DGYN RAKAYIATGGPLKSTFEDFWRMIWE DGYN RAKAYIATGGPLFOFFWRMIWE DGYN BKNIKTIATGGPLFOFFWRMIWE DGYN RAKAYIATGGPLFOFFWRMIWE DGYN RAKAYIATGGFLFOFFWRMIWE DGYN RAKAYIATGGFLFOFFWRMIWE DGYN RAKAYIATGGFLFOFFWRMIWE DGYN RAKAYIATGGFLFOFFWRMIWE DGYN RAKAYIATGGFLFOFFWRMIWE DGYN RAKAYIAT	PTP1B DyINASL1 DyINASL1 DiEEAQRSYILTGGPLPNTCGHFWEMVWE PTP_zela_D1 DGKLTDYINANYV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PTP_gamma_D PGKN RAKAYIAAGGPLKSTAEDFWRMIWE PTP_MUD1 PGS DYINANYI PTP_MUD1 PGS DYINANYI PTP_alpha_D1 PCS DYINANYI PTP_alpha_D1 PCS DYINANYI PCS DYINASYI DGYR KONAYIAAGGPLPFTFDCFWRMIWE DGYR KONAYIAAGGPLPFTFDCFWRMIWE DGYR RONAYIAAGGPLPFTFDCFWRMIWE DGYR RONAYIAAGGPLPFTFDCFWRMIWE DGYR RONAYIAAGGPLPFTFDCFWRMIWE DGYR RONAYIAAGGPLPFTFDCFWRMIWE DGYR RONAYIAAGGPLPFTFDCFWRMIWE DGYR RONAYIAAGGPLPFTFDFWRMIWE DGYR BPRKYIAAAGGPRGTTVDFWRMIWE DGYR BPRKYIAAAGGPRGTTVDFWRMIWE PCS DYINASYI PGBN FRREYIVTGGPLPGTKDDFWRMIWE PTP_DO1 PGS DYINANYIKNOLL PGBN FRREYIVTGGPLPGTKDDFWRMIWE PTP_DO1 PGS DYINANYIKNOLL PGBN FRREYIVTGGPLPGTKDDFWRMIWE PTP_DO1 PGS DYINANYIKNOLL PGBN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PTP_DO1 PGS DYINANYIKNOLL PGBN PREFIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVTGGPLPGTKDDFWRMIWE PGNN FRREYIVE PGNN FRREY	PTP1B DyINASLI DyINASLI DYINASLI DYINASLI DGKLTDYINANY DGYN RPKAYILTGGPLPNTCCHFWLMVWQ PTP_zela_D1 DGKLTDYINANY DGYN RPKAYILTGGPLFKSTAEDFWRMIWE PTP_gamma_D PGS DYINANY DGYN RPKAYILTGGPLFKSTAEDFWRMIWE DGYN RPKAYILTGGPLFKSTAEDFWRMIWE DGYN RPKAYILTGGPLFKSTAEDFWRMIWE DGYN RPKAYILTGGPLFFFEFFRMIWE DGYN RPKAYILTGGPLFFFFEFFRMIWE DGYN RPKAYILTGGPLFFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFWMIWE DGYN RAMAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFFWMIWE DGYN RPKAYILTGGPLFFFFWMIWE DGYN RPKAYILTGGPLFFFFWMIWE DGYN RPKAYILTGGPLFFFFWMIWE DGYN RPKAYILTGGPLFFFFWMIWE DGYN RPKAYILTGGPLFFFWMIWE DGYN RPKAYILTGGPLFFFWMIWE DGYN RPKAYILTGGPLFFFWMIWE DGYN RPKAYILTGGPLFFFWMIWE DGYN RPKAYILTGGPLFFFWMIWE DGYN RPWAYILTGGPLFFWMIWE DGYN RPWAYILTGGPLFWMIWE DGYN RPWAYILTGG	PTP1B Dyinasli Pyinasli PTP_zela_D1 DGKLTDyinanyv DGYN RPKAYIAGGPLENTCCHFWLMWW PTP_gamma_D DGKLTDyinanyv DGYN RAKAYIATGGPLENTCCHFWLMWW DGYN RAKAYIATGGPLESTAEDFWRMIWE PTP_gamma_D PGS Dyinanyi DGYN KAKAYIATGGPLESTFEDFWRMIWE DGYN KAKAYIATGGPLESTFEDFWRMIWE DGYN KAKAYIATGGPLESTFEDFWRMIWE DGYN KAKAYIATGGPLESTFEDFWRMIWE DGYN KAKAYIATGGPLESTFEDFWRMIWE DGYN KAKAYIATGGPLEDTFECFWRMIWE DGYN KAKAYIATGGPLEPFWRMIWE DGYN KAKAYIATGGPLEPFWRMIWE DGYN KAKAYIATGGPLEPFWRMIWE DGYN KONAYIATGGPLEPFWRMIWE DGYN KONAYIATGGPLEPFWRMIWE DGYN ENKKEIAAGGPECTVNDFWRMIWE DGYN ENKKEIAAGGPRETYNDFWRMIWE SH.PTP2 SH.PTP2 SH.PTP2 BPO SDYINANYIKNQLL GPDE NAKTYIAAGGPLEATVNDFWRWW PGLW SPREKYIATGGPLEATVNDFWRWW PGLW SPREKYIATGGPLEATVNDFWRW PGLW SPREKYIATGGPLEATVNDFWRW PGLW SPREKYIATGGPLEATVNDFWRW PGLW SPREKY	PTP1B Dyimasi Dyima	TCPTP DyINASLI DIEBAQRSYILITGGPLPNTCGHFWENVWE PTP_xi_D1 DGKLTDYINANY DGYN RPKAYILAGGPLRSTAEDFWRMIWE PTP_zela_D1 DGKLTDYINANY DGYN RPKAYIAAGGPLKSTAEDFWRMIWE PTP_gamma_D PKKNL DYINANY DGYN KAKAYIAAGGPLKSTAEDFWRMIWE PTP_gamma_D KKNL DYINANY DGYN KAKAYIAAGGPLKSTEDFWRMIWE PTP_alpha_D1 PGS DYINANYI PTP_alpha_D1 PDS DYINANYI PTP_alpha_D1 PDS DYINASYI DGYK EKNKFIAAGGPCETVNDFWRMIWE PTP_popsilon_D PCS DYINASYI DGYK EKNKFIAAGGPCETVNDFWRMIWE SH. PTP2 EPP_00a SH. PTP2 DYINASYI DGYK EKNKFIAAGGPCETVNDFWRMIWE SH. PTP2 PTP_00a SH. PTP2 DYINASYI PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PCS DYINANYIKNQLL PGNN PRREXIVTGGPLPTTCDFWRMIWE PTP_bola PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PTP_bola PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PTP_STEP PTP_D10 PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PTP_STEP PTP_D10 PCS DYINANYIKNQLL PGNN FRREXIVTGGPLPTTCDFWRMIWE PTP_STEP PTP_STEP PTP_STEP PGN SPQFFLATGGPLPTTCDFWRMIWE PTP_STEP PTP_STEP PTP_STEP PTP_STEP PGN SPQFFLATGGPLPTTRDFWRMIWE PTP_STEP PTP_STEP PTP_STEP PTP_STEP PGN SPQFFLATGGPLPTTRDFWRMIWE PGN SPQFFLATGGPLPTTRDFWRMIWE PGN SPQFFLATGGPLPTTRDFWRMIWE PGN SPGFLATGGPLPTTRDFWRMIWE PGN SPQFFLATGGPLPTTRDFWRMIWE PGN SPGFLATGGPLPTTRDFWRMIWE PGN SPGFLATGGPLPTTR	PTP1B DyINASLI DyINASLI DYINASLI DYINASLY DGKLTDDINANYV DGYN RPKAYILAGGPLPNTCCHFWLMVW DGYN RPKAYILAGGPLENTCCHFWLMVW DGYN RPKAYILAGGPLENTCCHFWLMW DGYN RACAYILAGGPLENTCCHFWLMW DGYN RACAYILAGGPLENTCCHFWLMW DGYN KAKAYILAGGPLENTREDFWRMIWE DGYN KAKAYILAGGPLEATVOGFWRWW DGYN BYNNANIWE DGYN BYNANIWE DGYN BYNNANIWE DGYN BYNNANIWE DGYN BYNNANIWE	PTP1B DyIMASIV DYIMASIV DGKLTDYIMANIV PTP_xi_D1 DGKLTDYIMANIV DGYN RPKAYIAAGGPLKSTAEDFWRMIWE PTP_gamma_D PGKLTDYIMANYI DGYN RAKAYIATGGPLETFEDFWRMIWE DGYN KAKAYIATGGPLETFEDFWRMIWE DGYN KAKAYIATGGPLETFEDFWRMIWE DGYN KAKAYIATGGPLETFEDFWRMIWE DGYN KAKAYIATGGPLETFEDFWRMIWE DGYN KAKAYIATGGPLETFRGDFWRMIWE DGYN KAKAYIATGGPLETTRGDFWRMIWE DGYN KAKAYIATGGPLETTRGDFWRMIWE PGN SDYIMANYI PGN SDYIMANYI PGN SPORENYIATGGPLETTRGDFWRMIWE PTP_STEP DYIMANYI PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PTP_PREST DYIMANYI PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PTP_PREST DYIMANYI PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SSYIMANYI RGYNGEBKVYIATGGPLETTRGDFWRMIWE PGN SSYIMANYI RGYNGEBKVYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PTP-PEPSAL DYIMASFW PGN SPORENYIATGGPLETTRGDFWRMIWE PTP-PEPSAL PTP-PEPSAL PDP SPORENYIATGGPLETTRGDFWRMIWE PTP-PEPSAL PTP-PEPSAL PTP-PEPSAL PREST PRESTIVTGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PTP-PEPSAL PRESTIVTGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PGN SPORENYIATGGPLETTRGDFWRMIWE PRESTIVETRGGPLETRGGPLETTRGGPLETRGGPLETTRGGPLETTRGGPLETTRGGPLETRGGPLETTRGGPLETRGGPLETRGGPLETRGGPL	PTP1B DyIMASLI DyIMASLI DyIMASLY DIEBAQRSYILTGGPLPNTCCHFWEMVWE DYIMASLY DGYN RPKAYIAAGGPLKSTAEDFWRMIWE PTP_Zela_D1 DGKLIDDIANNY PTP_Gamma_D PTP_Gamma_D PTP_Gamma_D PTP_ALPha_D1 PGS DYIMANYI PTP_DPS D1 TMS DYIMANYI PTP_DPS D1 TMS DYIMANYI PTP_DPS D1 TMS DYIMANYI PTP_DPS D1 TMS DYIMANYI PTP_DPS D2 TMS DYIMANYI PTP_DPS D2 TMS DYIMANYI PTP_DPS D3 TMS DYIMANYI PTP_DPS D4 TMS DYIMANYI PTP_DPS D4 TMS DYIMANYI PTP_DPS D4 TMS DYIMANYI PTP_DPS D5 TMS DYIMANYI PTP_DPS D5 TMS DYIMANYI PTP_DPS D5 TMS DYIMANYI PTP_DPS D5 TMS DYIMANYI PTP_STEP PTP D6 TMS DYIMANYI PTP D7 TMS DYIMANYI PTP D6 TMS PREFINTIAGPLPOTYUDFWRWWE PTP D7 TMS DYIMANYI TMS DYIMANYI PTP D7 TMS DYIMANYI TMS DAYNYI TMS DYIMANYI TMS DYIMANYI TMS DYIMANYI TMS DYIMANYI TMS DAYNYI TMS	PTP1B	PURPLE	POPINASLI POINASLI POINASLI POINASLI POINASLI DIEBAQRIIIJGGPLPNICCHFWEMVWE DIEBAQRSIIIJGGPLPNICCHFWEMVWE DIEBAQRSIIIJGGPLPNICCHFWEMVWE DIEBAQRSIIIJGGPLPNICCHFWEMWWE DIEBAQRSIIIJGGPLPNICCHFWEMWWE DIEBAQRSIIIJGGPLPNICCHFWEMWWE DIEBAQRSIIIJGGPLPNICCHFWEMWWE DIEBAQRSIIIJGGPLPNICCHFWEMWWE DIEBAGNETIANITH DIEBA	Publish	PUPPLE DYIMASLI KWEERAGRSYLITGGFLPWTCGHFWEMVWE DYIMASLY DIEEAGRSYLITGGFLPWTCGHFWEMVWE DYIMASLY DGYN RPKAYIAAGGFLKSTAEDFWAMIWE DYIMASLY DGYN RPKAYIAAGGFLKSTAEDFWAMIWE DGYN RPKAYIAAGGFLKSTAEDFWAMIWE DGYN RYAAXIAATGGFLKSTAEDFWAMIWE DGYN RACAYIAATGGFLKSTAEDFWAMIWE DGYN RACAYIAATGGFLKSTAEDFWAMIWE DGYN KACAYIATAGGFLEFDFWAMIWE DGYN KACAYIATAGGFLEFDFWAMIWE DGYN KACAYIATAGGFLEFDFWAMIWE DGYN RACAYIATAGGFLEFDFWAMIWE DGYN RACAYIATAGGFLEFDFWAMIWE DGYN RACAYIATAGGFLEFDFWAMIWE DGYN RACAYIATAGGFLEFDFWAMIWE DGYN RACAYIATAGGFLEFDFWAMIWE DGYN RACAYIATAGGFLEFTCHWEMWE DGYN RACAYIATAGGFLEF	PTP1B DyINASLI DyINASLI DYINASLI DIEBAQRSYILIGGPLPNTCGHFWIAWWO DIENAGRATIANGGPLKSTABDFWRMIWE DYINANIV DGYN RPKAYILIGGPLPDTCCHFWIAWWO DGYN RPKAYILIGGPLESTABDFWRMIWE PTP-94mma_D DSKHSDYLWANYV DGYN RRKAYINAGPLKSTABDFWRMIWE DGYN RYAYINAGPLKSTABDFWRMIWE DGYN RYAYINAGPLKSTABDFWRMIWE DGYN RYAYINATGPLEDTROCHFWIWE DAYN SPREIVITGGPLEATVODFWRMIWE DAYN SPREIVITGGPLEATVODFWRWIWE DAYN SPREIVITGGPLEATVODFWRWIWE DAYN SYNWEI DOLL SSYLWANYI DGYN RYAYINATGPLENTROCHFWIWWE DGYN RYAYINATGPLENTROCHFWIWWE DGYN RYAYINATGPLENTROCHFWIWWE DGYN RYAYINATGPLENTROCHFWIWWE DAYN RYAYINATGPLENTROCHFWIWWE DYINANYI DGAN RYAYINATGPLENTROCHFWIWWE DAYN RYAYINATGPLENTROCHFWIWWE DAYNWEI DOLL SSYLWANYI DOAN RYAYINATGPLENTROCHFWIWWE DAYNWEI DOAN RYGOT GGYNGTCAGFERINATE DYINASFI DOAN RYGOT GGYNGTCAGFERINATE DYINASFI DOAN RYCHATGGPLENTTALGFREWWEN DOAN RYCHATGGPLENTTALGFREWWEN DOAN RYCHATGGPLENTROCHFWIWWE DOAN RYCHATGGPLENTROCHFWIWWE DAYNWEI DOAN RYCHATGGPLENTROCHFWIWWE DAYNWEI DOAN RYCHATGGPLENTROCHFWIWWE DOAN RYCHATGGPLENTROCHFWIWWE DOAN RYCHATGGPLENTROCHFWIWWE DAYNWEI DOAN RYCHATGGPLENTROCHFWIWWE DOAN RYCHATGGPLENTROCHFWIWM RYCHATG	Public DyINASLI	PURBAQRSYILINGGPLPNTCGHFWEAWWE DyINASIN DyINASIN DHEAQRSYILINGGPLPNTCGHFWEAWWE DyINASIN DGYN RPKAYIAAGPLKSTABDFWEAUWE DYINASIN DGYN RPKAYIAAGPLKSTABDFWEAUWE DYRASIN DGYN RPKAYIAAGPLKSTABDFWEAUWE DGYN RAKAYIAAGPLKSTABDFWEAUWE DGYN RAKAYIATGGPLKSTABDFWEAUWE DGYN RAKAYIATGGPLKSTABDFWEAUWE DGYN RAKAYIATGGPLKSTBDFWEAUWE DGYN RAKAYIATGGPLETWADFWEAUWE DGYN RAKAYIATGGPLETWADFWEAUWE DGYN RAKAYIATGGPLETWADFWEAUWE DGYN RAKAYIATGGPLETWADFWEAUWE DGYN RAKAYIATGGPLETWADFWEAUWE DGYN RAKAYIATGGPLETWADFWEAUWE DGYN RAKHAYIATGGPLETWADFWEAUWE DGYN RAKHAYIATGGPLETWADFWEAUWE DGYN RAKHAYIATGGPLETWADFWEAUWE DGYN RAKHAYIA DGYN RAKHAWE DGYN BYLANAYI RAYAI BGYN RAKHAYIATGGPLANAYIA DGYN RAKHAYIA DGYN RAKHAYIA DGYN RAKHAYIA DGYN RAKHAYA DFWEAUWE DAYINANYI RAYAI RCYN GRENATAGGPLEATWADFWEAUWE DYN BYLANAYI RAYAI BCYN GRENATAGGPLEATWADFWEAUWE DYN RASY BYLANAYI BCYN GRAYATGGPLEATWADFWEAUWE DYN RASAL DYN BYLANAYI BCYN RAYAYATGGPLEATWADFWEAUWE DYN RASAL DYN RAYAYATGGPLEATWADFWEAUWE DYN RAYAYANAY BOYN REPRAITAGGPLEATWADFWEAUWE DYN RAYAYATGGPLEATWADFWEAUWE DYN RAYAYAYATGGPLEATWADFWEAUWE DYN RAYAYAYATGGPLEATWADFWEAUWE DYN RAYAYAYATGGPLEATWADFWEAUWE DYN RAYAYAYATGGPLEATWADFWEAUWE DYN RAYAYAYATGGPLEATWADFWEAUWE DYN RAYAYAYATGGPLEATWADFWEAUWE DYN RA	PURBAGE PAINASIN	PTP1B	PTP18	PTP1B DYINASLI DYINASLI DIEBAÇRSYILIAGPLPNTCGHFWENWE DYINASLY DGKLIDJIANNY DGKLIDJIANNY DGKLIDJIANNY DGKN RRYAYLAGGPLKSTREDFWENIWE PTP_Gamma_D DKKIDJIANNY DGKN RRYAYLAGGPLKSTREDFWENIWE PTP_GADI TNS DYINANY DGYN RRYAYLAGGPLKSTREDFWENIWE DGKN RAKAYLAGGPLKSTREDFWENIWE DGYN RAKAYLAGGPLKSTREDFWENIWE DGYN RAKAYLAGGPLKSTREDFWENIWE DGYN RAKAYLAGGPLKSTREDFWENIWE DGYN RAKAYLAGGPLKSTREDFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGREGATIONFWENIWE DGYN ROWAYLAGGREGATIONFWENIWE DGYN ROWAYLAGGREGATIONDFWENIWE DGYN ROWAYLAGGREGATIONDFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE PTP-DOI EGS DYINAST DYLNAST DYLNAST DOYN SPRETTYGGPLHGTRODFWENIWE DGYN ROWAYLAGGPLEFTWODFWENIWE DOYN SPRETTYGGPLHGTRODFWENIWE DOYN SPRETTYGGPLHGTRODFWENIWE DOYN SPRETTYGGPLHGTRODFWENIWE DOYN SPRETTYGGPLHGTRODFWENIWE DOYN SPRETTYGGPLHATADFWENIWE DYLNAST DOYN ROWAYLAGGPLEFTADFWENIWE DOYN REPAIN DOWN THE DOWN ROWAYLAGGPLEFTADFWENIWE DOYN ROWAYLAGGPLE	THE DYINASLY DIEBAGESTILGGELFREAWWR OKSECHFREAWWR OKSECHVALINEW PTP_2ela_D1 DRIEBAGESTILGGELFREATHER GRYGAVILAGEPLESTREDFREAKING OKTATAVATIANLY PTP_2ela_D1 DSKHEDFLANAYY DGYN RPKAYILAGGPLKSTREDFREAKING GYTKAVAVALINEW PTP_2ela_D1 DSKHEDFLANAYY DGYN RPKAYILAGGPLKSTREDFREAKING GYTKAVAVATIANLY DGYN RPKAYILAGGPLKSTREDFREAKING GYTKAVAVATIANLY DGYN RAPEA DGYN KKAYLAGGPLKSTREDFREAKING GYTATAVATIANLY DGYN KGANILAGGPLESTRODFREAKING GYTATAVATIANLY DGYN KGANILAGGPLESTRODFREAKING GYTATAVATIANLY DGYN KGANILAGGPLESTRODFREAKING GYTATAVATIANLY DGYN KGANILAGGPLESTRODFREAKING GYTATAVATIANLY BYTA DGYN KKAYLAGGPLESTRODFREAKING GYTATAVATIANLY BYTATAGGPLESTRODFREAKING GYTATAVATIANLY BYTATAGGPLESTRODFREAKING GYTATAVATIANLY GYTA BYTATAGGPLESTRODFREAKING GYTATAVATIANLY GYTA BYTATAGGPLESTRODFREAKING GYTATAVATIANLY GYTA BYTATAGGPLESTRODFREAKING GYTATAVATIANCH GYTA GYTA GYTATAGGPLESTRODFREAKING GYTATAVATIANCH GYTA GYTA GYTA GYTATAGGPLESTRODFREAKING GYTATATAGGPLESTRODFREAKING GYTATAGGPLESTRODFREAKING GYTATATAGGPLESTRODFREAKING GYTATAGGPLESTRODFREAKING GYTATAGGPLESTRODFREAKING GYTATAGGPLESTRODFREAKING GYTATAGGPLESTRODFREAKING GYTATAGGPLESTRO

		30 40 50	60
11: DTD1D	DEDCRUBKI, DKNKN	RYRDVSPEDHSRIKLHOE	NG
Kin TOTP	DYPHRVAKEPENRN	RNRYRDVSPYDHSRV LQNA	EN
Hum PTP xi D1	GITADSSNHPDNKH	KNRYINIVAYDHSRVKLAQL	AEK
PTP zel	GITADSSNHPDNKH.	KNRYINIVAYDHSRVKLAQL	AEK
Hum PTP gamma D	NITAEHSNHPENKH	KNRYINILAYDHSRVKIRPL	PGK
Dros PTP99A D1	DLPCEHSQHPENKR	KNRYLNITAYDHSRVHLHPT	PGO
Hum LCA D1	OFTWENSNLEVNKP	KNRYANVIAYDHSRVILTSI	DGV
Hum PTP mu D1	SAPWDSAKKDENRM	KNRYGNIIAYDHSRVRLQTI	EGD
Him PTP alpha D1	OATCEAASKEENKE	KNRYVNILPYDHSRVHLTPV	EGV
Hum PTP obsilon D	OGTFELANKEENRE	KNRYPNILPNDHSRVI <i>L</i> SQL	DGI
Mouso CD45 D1	KFPIKDARKPHNON	KNRYVDILPYDYNRVELSEI	NGD
Hum SH. PTP2	LYSRKEGOROENKN	KNRYKNILPFDHTRVVLHDG	DPN
Hum SH. PTP1	LHORLEGORPENKG	KNRYKNILPFDHSRVI <i>L</i> QGR	DSN
Hum PTP bola	NOSCDIALLPENRG	KNRYNNILPYDATRVKLSNV	aga
Dros PTP10D	DOPCTFADLPCNRP	KNRFTNILPYDHSRFKLQPV	ססס
Hum SAP. 1	SOSOMVASASENNA	KNRYRNVLPYDWSRVPLKPI	HEE
Ral PTP STEP	FVDPKEYDIPGLVR	KNRXKTILPNPHSRVRLTSP	DPE
Dros PTP69A D1	DRITKNSDLKENAC	KNRYPDIKAYDQTRVKLAVI	NGL
Hum MEG2	VGTFHCSMSPGNLE	KNRYGDVPCLDQTRVKLTKR	SGH
Hum PTP.PEST	IYPTATGEKEENVK	KNRYKDILPFDHSRVKLTLK	TPS
Hum PTPH1	GLAITFAKLPQNLD	KNRYKDVLPYDTTRVLLQGN	EDY
Dici PTP1	PSETSEGDKKHNTS	KNRYTNILPVNHT <i>R</i> VQ <i>L</i> KKI	ADK E
Fiss yeast_pyp1	QWSTVDSLSNTSYK	KNRYTDIVPYNCTRVHLKRT	SPS
Figs yeast_pyp2	WCCLASSRSTSISR	KNRYTDIVPYDKTRVRLAVP	KGC
Hum PTP xi D2	GITADSSNHPDNKH	KNRYINIVAYDHSRVKLAQL	AEK
Hum LCA D2	TSRFISANLPCNKF	KNRLVNIMPYELTRVCLQPI	RGV
Hum PTP alpha D2	NDKMRTGNLPANMK	KNRVLQIIPYEFNRVIIPVK	RGE
Hum PTP opsilon D2	KENMRTGNLPANMK	KARVIQIIPYDFNRVILSMK	RGQ
Hum PTP mu D2	VEDCSIALLPRNHE	KNRCMDILPPDRCLPFLITI	DGE
Mouse_CD45_D2	WRTQHIGNQEENKK	KNRNSNVVPYDFNRVPLKHELEMSKESEPESDESSDDDSD	SDDDSD
Dros PTP69A D2	SKSCSVGENEENINM	KNRSQEIIPYDRNRVILTPL	PMR
Hum PTP zeia D2	QSDYSAALKQCNRE	KNRTSSIIPVERSRVGISSL	SGE
Hum_PTP_gamma_D2	VECFSAQKECNKE	KNRNSSVVPSERARVGLAPL	PGM
Dros_PTP99A_D2	ETINIMABOVEBLKNCT	ETINIMAEOVEELKINCTPYLEOOXKINI IOFOPKO IHTASAMKOVINSI KINGALFPI EGSRVHI	PIEGSRVHLTPKP
Yarsinia_PTP	TNDPRYLOACGGEKI	CROTAVRA	
PTP1Bseq.no.	30 40	20 60	
PTP1B66			
	3	AAAA Figure 1B	

Figure 1B

5,6	
NLTTQETREI NINSGETRTI SQKGRPSGRVV SQKGRPSGRVV KKQCNTEKLV KKGCNTEKLV KKGCNTEKLI DMTNRKPORLI PDGCKAPRLV KKEKATGREV VGQGNTERTV LDNGDLIREI EEQLDAHRLI RGSEQRIL QVBEQKTLSV RGTEERGL VGBEEDRRQI NTGTEERTV LTFEGETRDI NTQTGEEHTV LTFEGETRDI NANFPSVKKV DKRNGSRRI SQKGRPSGRVV DARDGQSRTI NTRENKSRQI ARQEGVRVV ATQDDYVLEV SIODDYELTV REAGOKTISV	I
150 WINDEREREM IFEDTINIKATLISEDIKSYTTVLELE WINSGERTRI NINSGERTRI NINSGERTRI NINSGERTRI NINSGERTRI SQUERESGRAV WAP ADGSE EYGN FLYTQKSYQVLAYTYTFILRITKIKG SQKGRESGRAV WAP ADGSE EYGN IVYLINSTKIHACTYVESIRATKIKG SQKGRESGRAV WAP ADGSE IVYLINSTKIHACTYVESIRATKIKG SQKGRESGRAV WAP ADGSE IVYLINSTKIHACTYVESIRATKIKG SQKGRESGRAV WAP ADGSE IVYLINSTKIHACTYVESIRATKIKG SQKGRESGRAV WAP ARGTE IVYL IQVTLIDTVELATYTVALH KKGVGRIFTERI NYGORAPERI NYGO	88
150 EEKEM DDQEM DDQEM LFKETG ADGSE ADGSE EYGN ADGSE EYGN TENSE EYGN TYGN DQGCW TYGN DQGCW TYGN DQGCW TYGN DQGCW TYGN TYGN DQGCW TYGN TYGN TYGN TYGN TYGN TYGN TYGN TYGN	₩ Figure 1C
EKGSLKCA O EKGSLKCA O EKGRRKCD O EKGRRKCD O EKGRRKCD O EKGRRKCD O EKGRVKCC K EKGRVKCC O EKGRVKCC O EKGRVKCC O EKGRRKCD O EKGREKCD O ENGREKCC	1 a
Hum_PTP1B Hum_TCPTP Hum_PTP_xi_D1 Hum_PTP_gamma_D Dros_PTP99A_D1 Hum_LCA_D1 Hum_PTP_mu_D1 Hum_PTP_mu_D1 Hum_SH.PTP_D1 Hum_SH.PTP_1 Hum_SH.PTP_1 Hum_SAP.1 Ral_PTP_b0la Dros_PTP10D Hum_SAP.1 Ral_PTP_STEP Dros_PTP10D Hum_SAP.1 Ral_PTP_STEP Dros_PTP10D Hum_PTP_1 Dros_PTP11 Dici_PTP1 Fiss_yeast_pyp2 Fiss_yeast_pyp2 Fiss_yeast_pyp2 Hum_PTP_N1 Dici_PTP1 Dici_PTP1 Fiss_yeast_pyp2 Hum_PTP_alpha_D2 Hum_PTP_alpha_D2 Hum_PTP_alpha_D2 Hum_PTP_alpha_D2 Hum_PTP_alpha_D2 Hum_PTP_gamma_D2 Dros_PTP69A_D2 Hum_PTP_gamma_D2 Dros_PTP99A_D2 PTP1BSeq.no.	PTP1B66

250 260 270	PVVVHCSAgIGRSG								·									PILVHCSAGVGRTGTLV	VPSSAASLIDFLRVVRNQQSLAVSNMGARSKGQCPEPPIVVHCSAGIGKTGTFC	PICIHCSAGCGRIGAIC		RNV	NTIVHCSAGVGRIGIFI	PMFVHCSAqVGRTGTFI	X.				٠	MKYH KHASILVHCRDGSQQTGLFC				G PIVIVDRYGGAOACTFC	GSSAVADDSKLRPVIHCRAGVGRTAOLI	210 220		Organica Comment	
240	G VPESPASFLNFLFKVRES GS	ט	G VPEYSLPVLTFVRKAAYA	G VPEYSLPVLTFVRKAAYA	(0	G TPDHPLPVLNFVKKSSAA NPA	G VPEYPTPILAFLRRVKAC	G VPYHATGLLGFVROVKSK	G VPFTPIGMLKFLKKVKAC	G VPFTPIGMLKFLKKVKTL	G VPEDPHLLLKLRRRVNAF	G VPSDPGGVLDFLEEVHHK	G VPSEPGGVLSFLDQINQR	G VPETTQSLIQFVRTVRDY INRSP	G VPNPPQTLVRFVRAFRDR	ט	K TPDRAPPLLHLVREVEEAAQQEGPH	Σ	ෆ	YOFHYVNWPDH D VPSSFDSILDMISLMRKYQEHE DV	THLOYVAWPDH G IPDDSSDFLEFVNYVRSLRVDSE	z	HHYOYPNWSDC N SPENVKSMVEFLKYVNNSHGSG	×	ω Ω	U	U	G IPAEGKGMIDLIAAVOKO	RD TPVSKRSFLKLIRQVDKWQ		DGE	PKWPN PDSPISKTFELISVIKEEAANR D		KMI,HCPSwPEM SNPNSIYDFIVDVHERCNDY RNG	TAVSS	180 190 200			AA COMMAND
Ċ	HIND PTPIR LHFHYTTWPDF		5	<u>,</u> [2) -			num Oro alaha ni TOFHFISWPDF					Hum PTP bola RHFHYTVwPDH	Dros PTP10D RHFHFTTWPDF		TEP	ים ים	•	PEST	•	_	t pvol			•		um pro onailon D2 ROFHF		•		s c	, C	1			•	PTP1B66	PTP1B66

	Dac	200	310	320 330 340
Hum_PTP1B	LADTCLLLMDKR	KDPSSVDI	LLEMRKF	LIQTADQLRFSYLAVIEGAKFIMGD
Hum_TCPTP	LVDTCLVLMEKG		KQVLLINMRKY RMG	LIQTPDQLRFSYMAIIEGAKCIKGDSS
Hum_PTP_xi_D1	VLDSMLQQIQHE		FGFLKHIRSQRNY	LVQTEEQYVFIHDTLVEAILSKETEV
Hum_PTP_zela_D1	VLDSMLQQIQHE	_	FGFLKHIRSQRNY	LVQTEEQYVFIHDTLVEAILSKETEV
Hum_PTP_gamma_D	VIDSMLQQIKDK		LGFLKHIRTQRNY	LVQTEEQYIFIHDALLEAILGKETEV
Dros_PTP99A_D1	VLDAMLKQIQQK	•	FGFLRHI RAQRNF	LVQTEEQYIFLHDALVEALASGETNL
Hum_LCA_D1	VIDAMLERMKHE	KT VDI	YGHVTCMRSQRNY	MVQTEDQYVFIHEALLEAATCGHTEV
Hum_PTP_mu_D1	VIDIMLDMAERE	GV VDI	YNCVRELRSRRVN	MVQTEEQYVFIHDAILEACLCGDTSV
Hum PTP alpha D1	VIDAMLDMMHTE	RK VDV	YGFVSRIRAQRCQ	MVQTDMQYVFIYQALLEHYLYGDTEL
Hum PTP opsilon D	VIDAMMAMMHAE	VQV YQV	FEFVSRIKNORPO	MVQTDMQYTFIYQALLEYYLYGDTEL
Mouso_CD45_D1	GIDAMLEGLEAE	GK VDV	YGYVVKLRRQRCL	MVQVEAQYILIHQALVEYNQFGETEV
Hum_SH.PTP2	VIDILIDIIREK	GVDCDIDV	PKTIQMVRSQRSG	MVQTEAQYRFIYMAVQHYIETLQRRI
Hum_SH.PTP1	VIDMLMENISTK	GLDCDIDI	QKTIQMVRAQRSG	MVQTEAQYKFIYVAIAQFIETTKKKL
Hum PTP bola	ALDRILOQLDSK	ICV SC	YGAVHDLRLHRVH	MVQTECQXVYLHQCVRDVLRARKLRS
Dros_PTP10D	TLDRILQQINTS	ICV YC	FGIVYAMRKERVW	MVQTEQQYICIHQCLLAVLEGKENIVGP
Hum_SAP.1	ALDVLLRQLQSE	GL LGP	FSFVRKMRESRPL	MVQTEAQYVFLHQCICGSSNSQPRPQPR
Ral_PTP_STEP	ATSICCOOLRRE	GV VDI	LKTTCQLRQDRGG	MIQTCEQYQFVHHAMSLY
Dros_PTP69A_D1	ALDSLIQQLEEE	DS VSI	YNTVCDLRHQRNF	LVQSLKQYIFLYRALLDTGTFGNTDI
Hum MEG2	SLDICLAQLEEL	GT LINV	FOIVSRMRTORAF	SIQTPEQYYFCYKAILEFA
Hum_PTP.PEST	AIDYTWNLLKAG	KIPEEFNV		AVQTKEQYELVHRAIAQLFEKQLQLY
Hum PTPH1	TMETAMCLTERN	LP IYP	LDIVRKMRDQRAM	MVQTSSQYKFVCEAILRVY
Dici_PTP1	TAVIMMKKLDHYFKQLDYNSRIDFNL	LDYNSRIDFNL		MVQQLEQYLFCYKTILDEIYHRLNC
Fiss_yeast_pyp1	VLDTILRFPESKLSGFNPSVADSSDVV	FNPSVADSSDV	VFQLVDHI RKQRMK	MVQTFTQFKYVYD LIDSL
Fiss_yeast_pyp2	AVDOLLOVPKNILPK	TINLEDSKDFI	I FNCVNSLRSORMK	MVONFEOFKFLYD WVDYL
Hum_PTP_xi_D2	VLDSMLQQIQHE	GT VNI		
Hum LCA D2	TLSIVLERMRYE	GV VDM	FQTVKTLRTQRPA	•
Hum_PTP_alpha_D2	ALSTVLERVKAE	GI LDV	FQTVKSLRLQRPH	•
Hum_PTP_opsilon_D2	ALSNILERVKAE		FQAVKSLRLQRPH	•
Hum_PTP_mu_D2	AISIVCEMLRHQ		_	
Mouse_CD45_D2	ALFNLLESAETE			•
Dros_PTP69A_D2	AMCILVQHLRLE	KC VDI	CATTRKLRSQRTG	LINSYAQYEFLHRAIINY
Hum_PTP_zeia_D2	ALTTLMHQLEKE	VCCV SN	•	•
Hum_PTP_gamma_D2	ALTILSQQLENE	VCV AN	-	-
Dros_PTP99A_D2	AISSLAIEMEYC	ST ANV	YOYAKLYHNKRPG	VWTSSEDIRVIYN ILSFLPGNLNLLKR
Yarsinia_PTP	GAMCMNDSRNSO	NS'I	PDMVSOMRVORNG	MVOKDEOLDVLIK LAE
PTP1Bseq.no.	230 .	240	250	260 270 280
PTP1B66	İ	ب		Carrie Con voc.
	Po	Figure	1E a5	90

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Vmax and Km of 37kDa-PTP1B Mutants Toward RCML

Enzy	yme	(nmo	Vmax l/min/mg)	Km (nM)	Kcat (min ⁻¹)
wild	d typ	e	60200	102	2244
Tyr	46 -	S L	4120 4160	1700 1700	154 155
Glu	115	→ A → D	5700 5900	45 20	212 220
Lys	116	→ A	68600	150	2557
Lys	120	→ A	19000	80	708
Asp	181	→ A → E	0.61 97	≤126 10	0.023 3.6
His	214	→ A	700	20	26
Cys	215	→ S	0.026		0.00097
Arg	221	→ K → M	11 3.3	80 1060	0.41 0.12
Gln	262	→ A	720	9	27

Figure 2